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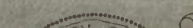
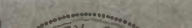
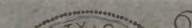
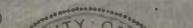
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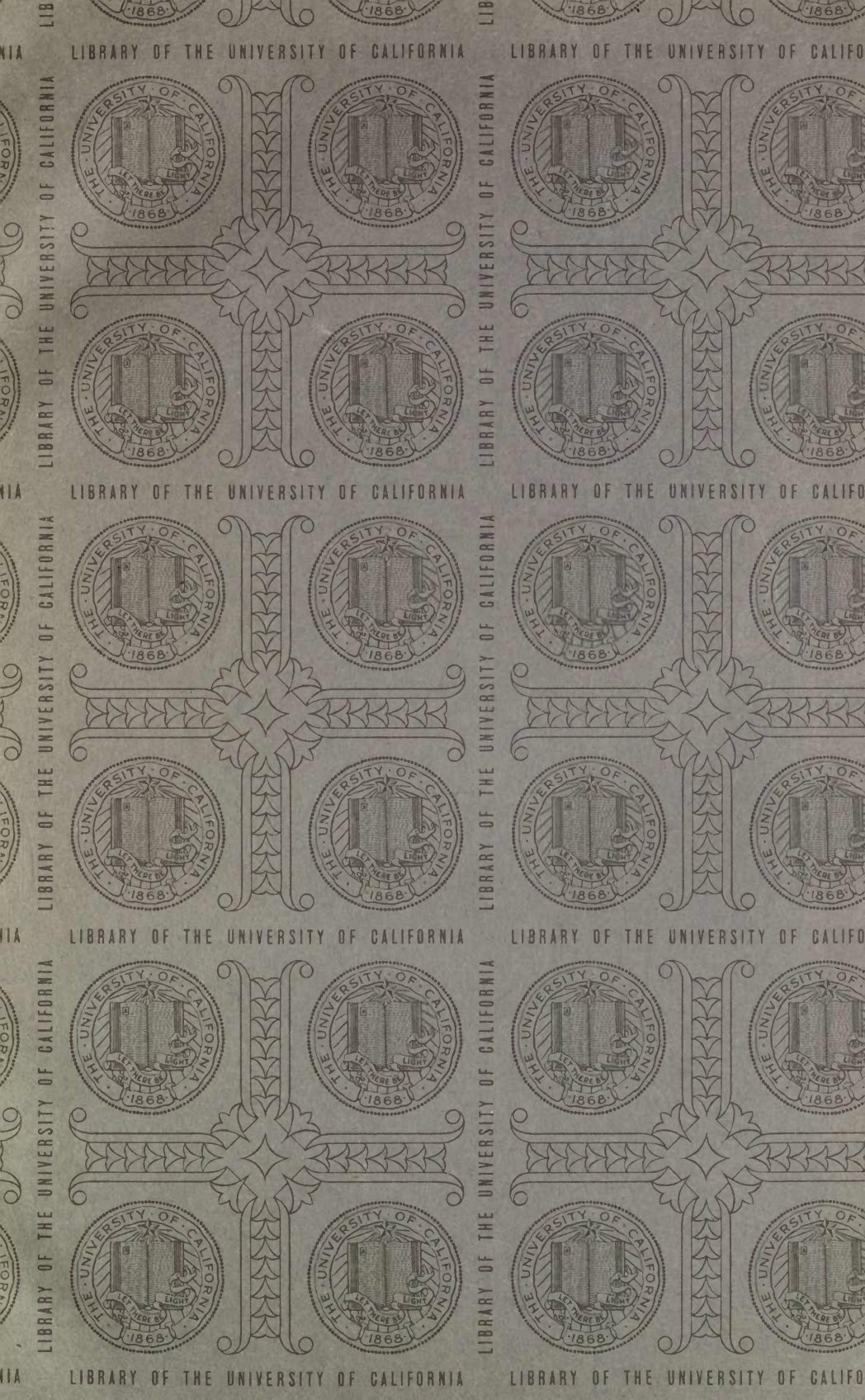
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TABLES OF
PHYSICAL AND
CHEMICAL CONSTANTS
AND SOME MATHEMATICAL FUNCTIONS

BY

G. W. C. KAYE

M.A., D.Sc., CAPT. R.E. (T.)

THE NATIONAL PHYSICAL LABORATORY

AND

T. H. LABY, M.A.

PROFESSOR OF NATURAL PHILOSOPHY, THE UNIVERSITY OF MELBOURNE

THIRD EDITION

LONGMANS, GREEN AND CO.

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1918

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PREFACE TO FIRST EDITION

THE need for a set of up-to-date English physical and chemical tables of convenient size and moderate price has repeatedly impressed us during our teaching and laboratory experience. We have accordingly attempted in this volume to collect the more reliable and recent determinations of some of the important physical and chemical constants.

To increase the utility of the book, we have inserted, in the case of many of the sections, a brief *résumé* containing references to such books and original papers as may profitably be consulted.

Every effort has been made to keep the material up to date ; in many cases a full reference to the original paper is given, while, failing such reference, the year of publication is almost always indicated.

The scope of the volume calls for little comment on our part. We have dipped a little into Astronomy, Engineering, and Geology in so far as they border on Physics and Chemistry. It will be noticed that considerable space has been allotted to Radioactivity and Gaseous Ionization : it is hoped that the collection of data, which we believe to be the first of the kind, will be of assistance to the numerous workers in a field whose phenomenal and somewhat transitional growth is a little dismaying from our present point of view.

Attention has been paid to the setting and accuracy of the mathematical tables ; these are included merely to facilitate calculations arising out of the use of the book, and limitations of space have cut out all but a few of the more essential functions. The convenience of the student of the newer physics has been studied by the inclusion of a table of values of e^{-x} reduced from Newman's original results.

We began this book while at the Cavendish Laboratory, Cambridge, and Dr. G. A. Carse shared in its inception. To Mr. G. F. C. Searle, F.R.S., we feel we owe much for his encouragement and suggestions when the scope of the book was under consideration. We record gratefully the help of a number of friends who have seen the proof-sheets of sections dealing with subjects with which their names are associated. Dr. J. A. Harker, F.R.S., and Mr. R. S. Whipple read the sections on Thermometry ; Mr. F. E. Smith revised the account of Electrical Standards, and Mr. C. C.

Paterson that of Photometry; Mr. A. Campbell criticized the section on Magnetism; and Professor Callendar, Principal Griffiths, and Dr. Chree have elucidated various points in Heat and Terrestrial Magnetism.

We owe thanks to Dr. Glazebrook for his permission to utilize the values of a number of constants recently determined at the National Physical Laboratory. Finally, we are greatly indebted to Mr. E. F. F. Kaye, M.Sc., who has given us valuable assistance in preparing the manuscript and revising the proof-sheets.

It was decided to keep the volume within reasonable limits, partly for the reader's convenience, and partly with the hope that the task of subjecting it to frequent revision in the future might not be impossible. We have consequently had to pick and choose our data, and it is scarcely likely that our selection will meet every individual requirement. That some sections are inadequately treated we fully realize, and we shall be very glad to receive suggestions and to be informed of any mistakes which, despite every care, have eluded us.

G. W. C. K.

T. H. L.

September, 1911.

PREFACE TO SECOND EDITION

WE regret that the difficulties of the times have not permitted the complete revision which we had contemplated. We have had to content ourselves with removing those mistakes of which, by the courtesy of many readers, we had become aware, and inserting a number of the more fundamental constants which contemporary research has yielded since 1911. A few tables have been thoroughly revised.

G. W. C. K.

T. H. L.

September, 1916.

PREFACE TO THIRD EDITION

IN the few months that have elapsed since the publication of the last edition, we have not found it possible to do more than bring a few primary constants up to date.

G. W. C. K.

T. H. L.

December, 1917.

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INTERNATIONAL ATOMIC WEIGHTS FOR 1918 (O = 16)

(See F. W. Clarke, "A Recalculation of the Atomic Weights," 1910)

Element.	Symbol.	Atomic Weight.	Element.	Symbol.	Atomic Weight.
Aluminium . . .	Al	27.1	Neodymium . . .	Nd	144.3
Antimony . . .	Sb	120.2	Neon	Ne	20.2
Argon	A	39.88	Nickel	Ni	58.68
Arsenic	As	74.96	Niobium† . . .	Nb	93.1
Barium	Ba	137.37	Niton (Ra. Em.)	Nt	222.0
Beryllium* . . .	Be	9.1	Nitrogen	N	14.01
Bismuth	Bi	208.0	Osmium	Os	190.9
Boron	B	11.0	Oxygen	O	16.00
Bromine	Br	79.92	Palladium	Pd	106.7
Cadmium	Cd	112.40	Phosphorus . . .	P	31.04
Cæsium	Cs	132.81	Platinum	Pt	195.2
Calcium	Ca	40.07	Potassium	K	39.10
Carbon	C	12.005	Praseodymium . .	Pr	140.9
Cerium	Ce	140.25	Radium	Ra	226.0
Chlorine	Cl	35.46	Rhodium	Rh	102.9
Chromium	Cr	52.0	Rubidium	Rb	85.45
Cobalt	Co	58.97	Ruthenium	Ru	101.7
Copper	Cu	63.57	Samarium	Sa	150.4
Dysprosium . . .	Dy	162.5	Scandium	Sc	44.1
Erbium	Er	167.7	Selenium	Se	79.2
Europium	Eu	152.0	Silicon	Si	28.3
Fluorine	F	19.0	Silver	Ag	107.88
Gadolinium . . .	Gd	157.3	Sodium	Na	23.00
Gallium	Ga	69.9	Strontium	Sr	87.63
Germanium	Ge	72.5	Sulphur	S	32.06
Gold	Au	197.2	Tantalum	Ta	181.5
Helium	He	4.00	Tellurium	Te	127.5
Holmium	Ho	163.5	Terbium	Tb	159.2
Hydrogen	H	1.008	Thallium	Tl	204.0
Indium	In	114.8	Thorium	Th	232.4
Iodine	I	126.92	Thulium	Tm	168.5
Iridium	Ir	193.1	Tin	Sn	118.7
Iron	Fe	55.84	Titanium	Ti	48.1
Krypton	Kr	82.92	Tungsten	W	184.0
Lanthanum	La	139.0	Uranium	U	238.2
Lead	Pb	207.20	Vanadium	V	51.0
Lithium	Li	6.94	Xenon	Xe	130.2
Lutecium	Lu	175.0	Ytterbium	Yb	173.5
Magnesium	Mg	24.32	Yttrium	Y	88.7
Manganese	Mn	54.93	Zinc	Zn	65.37
Mercury	Hg	200.6	Zirconium	Zr	90.6
Molybdenum . . .	Mo	96.0			

* Beryllium or Glucinum (Gl).

† Niobium or Columbium (Cb).

The following atomic weights for 1911 (see pp. 109, 127) include only those which have been subsequently changed:—

Ca, 40.09; C, 12.00; Er, 167.4; He, 3.99; Fe, 55.85; Kr, 82.9; Pb, 207.10; Lu, 174.0; Hg, 200.0; Pr, 140.6; Ra, 226.4; S, 32.07; Ta, 181.0; Th, 232.0; Sn, 119.0; U, 238.5; V, 51.06; Yb, 172.0; Y, 89.0.

THE ELEMENTS IN THE ORDER OF ATOMIC WEIGHTS (1918)

Symbol.	Atomic Weight.	First isolated by	Date.	Symbol.	Atomic Weight	First isolated by	Date.
H	1.008	Cavendish	1766	Ru	101.7	Claus	1845
He	4.00	Ramsay & Cleve *	1895	Rh	102.9	Wollaston	1803
Li	6.94	Arfvedson	1817	Pd	106.7	Wollaston	1803
Be§	9.1	Wöhler and Bussy	1828	Ag	107.88	—	P.
B	11.0	Gay-Lussac & Thénard	1808	Cd	112.40	Stromeyer	1817
C	12.005	—	P.	In	114.8	Reich and Richter	1863
N	14.01	Rutherford	1772	Sn	118.7	—	P.
O	16.00	Priestley and Scheele	1774	Sb	120.2	Basil Valentine	15 centy.
F	19.0	Moissan	1886	I	126.92	Courtois	1811
Ne	20.2	Ramsay and Travers	1898	Te	127.5	v. Reichenstein	1782
Na	23.00	Davy	1807	Xe	130.2	Ramsay and Travers	1898
Mg	24.32	Liebig and Bussy	1830	Cs	132.81	Bunsen and Kirchhoff	1861
Al	27.1	Wöhler	1827	Ba	137.37	Davy	1808
Si	28.3	Berzelius	1823	La	139.0	Mosander	1839
P	31.04	Brand	1674	Ce	140.25	Mosander	1839
S	32.06	—	P.	Pr	140.9	Auer von Welsbach	1885
Cl	35.46	Scheele	1774	Nd	144.3	Auer von Welsbach	1885
K	39.10	Davy	1807	Sa	150.4	L. de Boisbaudran	1879
A	39.88	Rayleigh & Ramsay	1894	Eu	152.0	Demarçay	1901
Ca	40.07	Davy	1808	Gd	157.3	Marignac	1886
Sc	44.1	Nilson and Cleve	1879	Tb	159.2	Mosander	1843
Ti	47.9	Gregor	1789	Dy	162.5	U. & D.	1907
V	51.0	Berzelius	1831	Ho	163.5	L. de Boisbaudran	1886
Cr	52.0	Vauquelin	1797	Er	167.7	Mosander	1843
Mn	54.93	Gahn	1774	Tm	168.5	Cleve	1879
Fe	55.84	—	P.	Yb	173.5	Marignac	1878
Ni	58.68	Cronstedt	1751	Lu	175.0	Urbain	1908
Co	58.97	Brand	1735	Ta	181.5	Eckeberg	1802
Cu	63.57	—	P.	W	184.0	Bros. d'Elhujar	1783
Zn	65.37	Ment. by B. Valentine	15 centy.	Os	190.9	Smithson Tennant	1804
Ga	69.9	L. de Boisbaudran	1875	Ir	193.1	Smithson Tennant	1804
Ge	72.5	Winkler	1886	Pt	195.2	—	16 centy.
As	74.96	Albertus Magnus	13 centy.	Au	197.2	—	P.
Se	79.2	Berzelius	1817	Hg	200.6	Md. by Theophrastus	300 B.C.
Br	79.92	Balard	1826	Tl	204.0	Crookes	1861
Kr	82.92	Ramsay and Travers	1898	Pb	207.20	Mentd. by Pliny	P.
Rb	85.45	Bunsen and Kirchhoff	1861	Bi	208.0	Mtd. by B. Valentine	15 centy.
Sr	87.63	Davy	1808	Nt	222	M. & Mme. Curie	1900
Y	88.7	Wöhler	1828	Ra	226.0	Curies and Bemont	1902
Zr	90.6	Berzelius	1825	Th	232.4	Berzelius	1828
Nb	93.1 †	Haichett	1801	U	238.2	Pelilot	1841
Mo	96.0	Hjelm	1790				

P., Prehistoric; * Lockyer (in-sun), 1868; U. & D., Urbain & Dementitroux; § Be or Ge; † Nb or Cb.

C.G.S. UNITS AND DIMENSIONS

References: Mach, "Science of Mechanics;" Everett, "C.G.S. System of Units;" Maxwell "Theory of Heat."

The metric standards of length and mass are kept at the International Bureau of Weights and Measures in the Pavillon de Breteuil, Sèvres, near Paris. The Bureau is jointly maintained by the principal civilized governments as members of the Metric Convention. The use of metric weights and measures was legalized in the United Kingdom in 1897.

LENGTH

Unit—the **centimetre**, $1/100$ of the international metre, which is the distance, at the melting-point of ice, between the centres of two lines engraved upon the polished "neutral web" surface of a platinum-iridium bar of a nearly X-shaped section, called the **International Prototype Metre**.

The alloy of 90 Pt, 10 Ir used (also for the International Kilogramme) has not a large expansion coefficient (see p. 53), is hard and durable, and was artificially aged. Pt-Ir copies of this metre, called **National Prototype Metres**, were made at the same time, and distributed by lot about 1889 to the different governments. The international metre is a copy of the original Borda platinum standard—the *mètre des archives*. This was intended to be one ten-millionth of the quadrant from the equator to the pole through Paris, and was legalized in 1795 by the French Republic. But as the value of a quadrant came to be more accurately determined, and moreover is changing, the actual bar constructed was made the standard.*

The international prototype metre has been measured (1894 and 1907) in terms of the wave-lengths of the cadmium rays (see p. 75), and equals $1,553,164.1$ wave-lengths of the red ray in dry air at 15° C. (H. Scale) and 760 mm. pressure. (See Michelson's "Light Waves," 1903.)

References: Guillaumé, "La Convention du Mètre," and Chree, *Phil. Mag.*, 1901.

MASS

Unit—the **gramme**, $1/1000$ of the **International Prototype Kilogramme**, which is the mass of a cylinder of platinum-iridium.

The international kilogramme is a copy of the original Borda platinum kilogramme—the *kilogramme des archives*—which was intended to have the same mass as that of a cubic decimetre of pure water at the temperature of its maximum density. More exact measurements revealed the incorrectness of the relation (see p. 10), and so the kilogramme was subsequently defined as above.

As with the metre, Pt-Ir copies of the international standard—**National Prototype Kilogrammes**—have been distributed to the different governments.

TIME

Unit—the **second**, which may be defined simply as $1/86,164.09$ of a **sidereal day**. For all practical purposes the sidereal day may be regarded as the period of a complete axial rotation (360°) of the earth with respect to the fixed stars.†

The second is usually defined as $1/(24 \times 60 \times 60)$ of a **mean solar day**, i.e. $1/86,400$ of the **average** value of the somewhat variable interval (the apparent solar day) between two successive returns of the sun to the meridian (see p. 15).

Strictly, the sidereal day is the interval between two successive transits of the first point of Aries‡ across any selected meridian.§ The true period of rotation of the earth is actually about $1/100$ second longer than the sidereal day; the difference arises from the slow and continual change of direction ("precession") of the earth's axis in space.

A **tropical or solar year** is the average interval between two successive returns of the sun to the first point of Aries; it is found to equal 365.2422 mean solar days. Our modern (Julian) calendar assumes that in 4 successive civil years, 3 consist of 365 days, and 1 of 366; the average thus being 365.25 days. The Gregorian correction (that century years are not to count as leap years unless divisible by 400) reduces this value to 365.2425 mean solar days, and thus the **average civil year** is a close approximation to a tropical year.

* According to the latest estimates, the *mean* meridian quadrant = $10,002,100$ metres (see p. 13).

† Tidal friction is retarding the rotation of the earth, so that the above (sidereal) definition of the second, while practically justified, is theoretically not quite perfect.

‡ The first point of Aries is that one of the two nodes of intersection of the ecliptic and the celestial equator where the sun (moving in the ecliptic) crosses the equator from south to north (at about March 21). The ecliptic is the apparent yearly track of the sun in a great circle on the celestial sphere.

§ Neglecting small irregularities, this is true also for any star.

BRITISH UNITS

A **sidereal year** is the time interval in which the sun appears to perform a complete revolution with reference to the fixed stars; *i.e.* it is the time in which the earth describes one sidereal revolution round the sun. Owing to precession, a sidereal year is longer than a tropical year.

	h.	m.	s.	
Mean solar day	= 24	0	0	= 86,400 secs.
Sidereal day	= 23	56	4 [·] 0906	= 86,164 [·] 0906 secs.
Tropical year	= 365 [·] 2422	mean solar days.		
Sidereal year	= 365 [·] 2564	"	"	" (epoch 1900).
	= 366 [·] 2564	sidereal days.		

Reference: Newcomb, "Astronomy."

BRITISH IMPERIAL STANDARDS.

(From information supplied by Major MacMahon, F.R.S., Board of Trade, Standards Office.)

According to the Weights and Measures Act, 1878, the **yard** is the distance, at 62° F., between the central transverse lines in two gold plugs in the bronze bar, called the **Imperial Standard Yard**, when supported on bronze rollers in such manner as best to avoid flexure of the bar.

The defining lines are situated at the bottom of each of two holes, so as to be in the median plane of the bar, which is of 1 inch square section and 38 inches long. Its composition is 32 Cu, 5 Sn, 2 Zn. Copper alloys are now known not to be suitable for standards of length, and in 1902 a Pt-Ir X-shaped copy of the yard was made.

The **pound** is the **weight** in vacuo of a platinum cylinder called the **imperial standard pound**.

The imperial standard yard and pound are preserved at the Standards Office of the Board of Trade, Old Palace Yard. A number of official copies have been prepared, and are in the custody of the Royal Society, the Mint, Greenwich Observatory, and the Houses of Parliament.

The **gallon** contains 10 lbs. weight of distilled water weighed in air against brass weights at a pressure of 30 inches, and with the water and the air at 62° F.

[NOTE.—No mention is made in the Act of the density of the brass weights, or of the humidity of the air.]

BRITISH AND METRIC EQUIVALENTS

The present legal equivalents are those legalized by the Order in Council of May 19, 1898, and derived at the International Bureau of Weights and Measures, by Benoit in 1895 in the case of the yard and the metre, and by Broch in 1883 for the pound and the kilogramme. (See *Trav. et Mém. du Bur. Intl.*, tomes iv, 1885, and xii, 1902.)

Imperial Standard.		International Prototype.	(Reciprocal.)
1 yard	=	·914399 metre	1 [·] 093614
1 pound	=	·45359243 kilogramme	2 [·] 2046223

[NOTE.—The yard is defined at 62° F., the metre at 0° C.]

DERIVED C.G.S. UNITS AND STANDARDS

GENERAL AND MECHANICAL UNITS

Area:—*Unit*—the square centimetre.

Volume:—*Unit*—the cubic centimetre (c.c.). The metric unit is the **litre**, now defined as the volume of a kilogramme of pure, air-free water at the temperature of maximum density (see p. 22) and 760 mm. pressure (*Procès Verbaux*, 1901, p. 175). The litre was originally intended to be 1 cubic decimetre or 1000 c.c.s.; the present accepted experimental relation is that 1 kilogramme of water at 4° C. and 760 mm. pressure measures 1000[·]027 c.c.s. (see p. 10).

Density:—*Unit*—grammes per c.c. **Specific gravity** expresses the density of a substance relative to that of water, and is objectionable in requiring two temperatures to be stated.

Velocity:—*Unit*—1 cm. per second. **Angular Velocity**:—*Units*—1 radian (57°·296) per sec.; 1 revolution per sec.

Acceleration:—Time rate of alteration of velocity. *Unit*—(1 cm. per sec.) per sec. **Angular Acceleration**:—*Units*—1 radian per sec.²; 1 revolution per sec.²

Momentum:—Mass multiplied by velocity. *Unit*—1 gm. cm. sec.⁻¹.

Moment of Momentum:—Momentum multiplied by distance from axis of reference. *Unit*—1 cm.² gm. sec.⁻¹.

Moment of Inertia:— Σmd^2 , where m is the mass of any particle of a body, and d its distance from the axis of reference. *Unit*—1 cm.² gm. (see p. 16).

Angular Momentum:—Moment of inertia multiplied by angular velocity round axis of reference. *Unit*—1 cm.² gm. sec.⁻¹.

Force:—Measured by the acceleration it produces in unit mass. *Unit*—the **dyne** = cm. gm./sec.² *Gravitational unit*—the weight of 1 gram = g dynes.

Couple, Torque, Turning Moment:—Force multiplied by distance from point of reference. *Unit*—1 dyne cm.

Work:—Force multiplied by distance through which point of application of force moves in direction of force. *Unit*—the **erg** = 1 dyne cm.; 1 **joule** = 10⁷ ergs. [1 calorie = 4·18 joules]. *Gravitational unit*—weight of 1 gm. \times 1 cm. = g dyne cms. = g ergs.

Energy:—Measured by the work a body can do by reason of either (1) its motion—**Kinetic Energy** ($= mv^2/2$) or (2) its position—**Potential Energy**. *Unit*—the **erg**. (See "Work.") 1 **Board of Trade Unit** = 1 kilowatt hour = $3\cdot6 \times 10^6$ watt-secs.

Power:—Work per unit time. *Unit*—1 erg per sec. 1 **watt** = 10⁷ ergs per sec. = 1 joule per sec. = 1 volt-ampere. 1 kilowatt = 1·34 horse-power.

Pressure, Stress:—Force per unit area. *Unit*—1 dyne per cm.² 1 **megabar** = 10⁶ dynes per cm.² = 750 *mm. mercury at 0° C., lat. 45°, and sea-level ($g = 980\cdot6$). 1 **atmosphere** = 760 mm. mercury at 0° C., lat. 45°, and sea-level = 759·4 mm. mercury at 0° C. in London = 1·0132 \times 10⁶ dynes per cm.² = 14·7 lbs. per inch² = 0·94 ton per foot².

* Correct to 1 part in 5000.

Elasticity:—Ratio of stress to resulting strain. *Unit*—1 dyne per cm.², since the dimensions of a strain are zero.

HEAT UNITS

Temperature:—The melting-point of pure ice under 1 atmosphere is defined as 0° C., and the boiling-point of water under 1 atmosphere as 100° C. This fundamental interval is divided into 100 parts by use of the constant-volume hydrogen thermometer (see p. 44); each part is a degree Centigrade. Dimensions of temperature are not required, as it is defined independently of mass, length, and time.

Heat:—*Dynamical unit*—the **erg**. *Thermal unit*—the **calorie** = heat required to raise the temperature of 1 gramme of water from t° C. to $(t+1)^\circ$ C. The **20° calorie** ($t = 20^\circ$) = 4·180 \times 10⁷ ergs. The **15° calorie** ($t = 15^\circ$) = 4·184 \times 10⁷ ergs. The **mean calorie** (= 1/100 heat required to raise 1 gramme of water from 0° to 100° C.) = 4·184 \times 10⁷ ergs. (see pp. 55, 56). 1 **watt-minute** = 14·3 calories. The large calorie = 1000 calories.

Gas Constant R , in $p v = R \theta / m$, where p is the pressure, v the volume, θ the absolute temperature of a gram-molecule (*i.e.* m grams) of a gas of molecular weight m . For 1 gram-molecule of an ideal gas of density ρ , $R = \frac{p v m}{\theta} = \frac{p}{\theta} \cdot \frac{m}{\rho} = \frac{1\cdot0132 \times 10^6 \times 22\cdot412}{273\cdot1} = 83\cdot15 \times 10^6$ ergs per gm. mol. (Berthelot,

see p. 106). This value is a constant for all ideal gases. To derive R for 1 gram of a gas, this figure should be divided by the molecular weight (oxygen = 16) of the gas. R has the dimensions of a specific heat in dynamical units.

ELECTRICAL AND MAGNETIC UNITS

Reference:—J. J. Thomson, "Mathematical Theory of Electricity and Magnetism." The fundamental basis of the electrostatic system of units is the repulsive force between two quantities of like electricity. In the electromagnetic system the repulsion between two like magnetic poles is taken as the basis.

The electromagnetic system (or one based on it) is universally employed in electrical engineering; the electrostatic is used only in certain special cases.

ELECTROSTATIC UNITS

Quantity or Charge:—*Unit*—that quantity which placed 1 cm. distance from an equal like quantity repels it with a force of 1 dyne.

ELECTRICAL UNITS

Current:—*Unit*—Unit quantity flowing uniformly past a point in unit time.

Potential Difference and Electromotive Force:—*Unit*—that P.D. which exists between two points when the work done in taking unit quantity from one point to the other is 1 erg.

Capacity:—*Unit*—the charge on a conductor which is at unit potential; or in the case of a condenser, when its plates are at unit P.D.

Dielectric Constant, Inductivity, or Specific Inductive Capacity of a medium is the ratio of the capacity of a condenser having the medium as dielectric, to the capacity of the same condenser with a vacuum as dielectric (p. 84).

ELECTROMAGNETIC UNITS

Magnetic Pole Strength or Quantity:—*Unit*—that quantity which, placed 1 cm. distance from an equal like quantity, repels it with a force of 1 dyne.

Magnetic Force or Field Strength:—*Unit*—the force which acts on unit magnetic pole.

Magnetic Moment of magnet = pole strength \times length of magnet.

Intensity of Magnetization = magnetic moment per unit volume.

Permeability of a medium is the ratio of the magnetic induction in the medium to that in the magnetizing field (p. 89).

Susceptibility:—*Unit*—intensity of magnetization per unit field (p. 89).

Electric Current:—*Unit*—that current which produces unit magnetic force at the centre of a circle of radius 2π cms.

Quantity = current \times time.

Potential and E.M.F.:—*Unit*—that P.D. which exists between two points when the work done in taking unit quantity from one point to the other is 1 erg.

Electrostatic Capacity = quantity/potential difference.

Resistance = potential difference/resulting current. (Ohm's law is assumed.)

Conductance:—Reciprocal of resistance.

Specific Resistance:—Resistance of prism of unit area and unit length.

Conductivity:—Reciprocal of specific resistance.

Coefficient of Self-induction of a circuit is the E.M.F. produced in it by unit time-rate of variation of the current through it.

Coefficient of Mutual Induction of two circuits is the E.M.F. produced in one by unit time-rate of variation of the current in the other.

PRACTICAL ELECTRICAL UNITS

At an International Conference on Electrical Units and Standards held in London, October, 1908, it was resolved that—

1. The magnitudes of the fundamental electrical units shall, as heretofore, be determined on the electromagnetic system of measurement with reference to the centimetre, gramme, and second (c.g.s.). These fundamental units are (1) the **Ohm**, the unit of electrical resistance, which has the value 10^9 c.g.s.; (2) the **Ampere**, the unit of electric current, which has the value 10^{-1} c.g.s.; (3) the **Volt**, the unit of electromotive force, which has the value 10^8 c.g.s.; (4) the **Watt**, the unit of power, which has the value 10^7 c.g.s. [For absolute electrical units, see p. 8.]

2. As a system of units representing the above, and sufficiently near to them to be adopted for the purpose of electrical measurements, and as a basis for legislation, the Conference recommends the adoption of the International Ohm, the International Ampere, and the International Volt.

3. The **Ohm** is the first primary unit. The **International Ohm** is defined as the resistance offered to an unvarying electric current by a column of mercury at 0° C., 14.4521 grammes in mass, of a constant cross-section, and of a length of 106.300 cms.

4. The **Ampere** is the second primary unit. The **International Ampere** is defined as the unvarying electric current which, when passed through a solution of nitrate of silver in water, in accordance with authorized specification, deposits silver at the rate of .00111800 gramme per second.

5. The **International Volt** is defined as the electrical pressure which, when steadily applied to a conductor whose resistance is one International Ohm, will produce a current of one International Ampere.

6. The **International Watt** is defined as the energy expended per second by an unvarying electric current of one International Ampere under an electric pressure of one International Volt.

DIMENSIONS OF UNITS

DIMENSIONS OF UNITS

The dimensions in terms of length, mass, and time are denoted by the indices given under L, M, and T. Thus the dimensions of power are L^2MT^{-3} .

MECHANICAL AND HEAT UNITS

Quantity.	L. M. T.	Quantity.	L. M. T.	Quantity.	L. M. T.
Length . . .	1 0 0	Momentum . . .	1 1 -1	Strain . . .	0 0 0
Mass . . .	0 1 0	Moment of mo-		Elasticity . . .	-1 1 -2
Time . . .	0 0 1	mentum . . .	2 1 -1	Compressibility	1 -1 2
Angle . . .	0 0 0	Moment of in-		Viscosity . . .	-1 1 -1
Surface . . .	2 0 0	ertia . . .	2 1 0	Diffusion . . .	2 0 -1
Volume . . .	3 0 0	Angular mo-		Capillarity . . .	0 1 -2
Density . . .	-3 1 0	mentum . . .	2 1 -1	Temperature . . .	0 0 0
Velocity . . .	1 0 -1	Force . . .	1 1 -2	Heat * . . .	2 1 -2
Angular vel. . .	0 0 -1	Couple, Torque	2 1 -2	Thermal Con	
Acceleration . .	1 0 -2	Work, Energy	2 1 -2	ductivity * . .	1 1 -3
Angular accele-		Power . . .	2 1 -3	Entropy * . .	2 1 -2
ration . . .	0 0 -2	Pressure, Stress	-1 1 -2		

ELECTRICAL AND MAGNETIC UNITS

v , the ratio of the electromagnetic to the electrostatic unit of quantity, is usually taken as 3×10^{10} , and is a pure number (p. 69). (See Rücker, *Phil. Mag.*, 22, 1889.)

Unit.	Sym- bol.	Dimensions.		Relations.			
		E.S. Unit.	E.M. Unit.	E.S.U.	Practical Unit.		
		L. M. T. k .	L. M. T. μ .	EMU.			
Electrical						E.M.U.	E.S.U.
Charge or quantity	e	$\frac{3}{2} \frac{1}{2} -1 \frac{1}{2}$	$\frac{1}{2} \frac{1}{2} 0 -\frac{1}{2}$	$1/v$	coulomb	$= 10^{-1}$	$= 3 \times 10^9$
Resistance	R	$-1 0 1 -1$	$-1 0 -1 -1$	v^2	ohm	$= 10^9$	$= \frac{1}{3} \times 10^{-11}$
Current	i	$\frac{3}{2} \frac{1}{2} -2 \frac{1}{2}$	$\frac{1}{2} \frac{1}{2} -1 -\frac{1}{2}$	$1/v$	ampere	$= 10^{-1}$	$= 3 \times 10^9$
Potential or E.M.F.	E	$\frac{1}{2} \frac{1}{2} -1 -\frac{1}{2}$	$\frac{3}{2} \frac{1}{2} -2 \frac{1}{2}$	v	volt	$= 10^8$	$= 1/300$
Electric field	F	$-1 \frac{1}{2} \frac{1}{2} -1 -\frac{1}{2}$	$\frac{1}{2} \frac{1}{2} -2 \frac{1}{2}$	v	(volt/cm.)	$=$	$=$
Conductivity	K	$0 0 -1 1$	$-2 0 1 -1$	$1/v^2$	"reciprocal ohm"	$= 10^{-9}$	$= 9 \times 10^{11}$
Capacity	C	$-1 0 0 1$	$-1 0 2 -1$	$1/v^2$	microfarad †	$= 10^{-15}$	$= 9 \times 10^5$
Self and mutual induction }	$L; M$	$-1 0 2 -1$	$1 0 0 1$	v^2	henry	$= 10^9$	$= \frac{1}{9} \times 10^{-11}$
Dielectric constant †	k	$0 0 0 1$	$-2 0 2 -1$	$1/v^2$	cm.	$= 1$	$= \frac{1}{3} \times 10^{-20}$
Magnetic							
Pole strength	m	$\frac{1}{2} \frac{1}{2} 0 -1$	$\frac{3}{2} \frac{1}{2} -1 \frac{1}{2}$	v	—	—	—
Flux (total lines)	N	$\frac{1}{2} \frac{1}{2} 0 -1$	$\frac{3}{2} \frac{1}{2} -1 \frac{1}{2}$	$1/v$	maxwell	$= 1$	$= 3 \times 10^{10}$
Force; field strength	H	$\frac{1}{2} \frac{1}{2} -2 \frac{1}{2}$	$-1 \frac{1}{2} \frac{1}{2} -1 -\frac{1}{2}$	$1/v$	gauss	$= 1$	$= 3 \times 10^{10}$
Induction	B	$-1 \frac{1}{2} \frac{1}{2} 0 -\frac{1}{2}$	$-2 \frac{1}{2} \frac{1}{2} -1 \frac{1}{2}$	v	gauss	$= 1$	$= \frac{1}{3} \times 10^{-10}$
Intensity of magnetization	I	$-3 \frac{1}{2} \frac{1}{2} 0 -\frac{1}{2}$	$-2 \frac{1}{2} \frac{1}{2} -1 \frac{1}{2}$	v	—	—	—
Permeability	μ	$-2 0 2 -1$	$0 0 0 1$	v^2	—	—	—

* In dynamical units.

† Specific inductive capacity.

‡ 10^{-6} farad.

Example:—To find the number (n) of ergs per sec. in a horse-power (33,000 ft.-lbs. per min.).

Dimensions of power = $L^2MT^{-3} = LT^{-1}$ [Force]

$$n = 33,000 \frac{\text{ft. (min.)}^{-1}}{\text{cm. (sec.)}} \cdot \frac{\text{lb. weight}}{\text{dyne}} = \frac{33,000 \times 30 \cdot 48}{60} \times 453 \cdot 6 \times 981$$

$$= 7 \cdot 46 \times 10^9 \text{ ergs per sec.} = 746 \text{ watts.}$$

ELECTRICAL UNITS

ABSOLUTE DETERMINATIONS OF ELECTRICAL UNITS

See Baillehache, "Unités Électriques," Paris, 1909, and the "Report of the London Conference" (p. 6). The appendix to this report (issued separately, 9d.) gives full particulars as to the realization of the ampere and ohm, together with the specification of the Weston normal (cadmium) cell.

THE OHM

The mean value **106.25** cms. of Hg of 1 sq. mm. cross-section at 0° C. may be taken as a measure of the present experimental value of the true ohm, which is equal to 10^9 E.M. (c.g.s.) units. Compare the international ohm (p. 6).

cm./0°.	Method.	Observer.	cm./0°.	Method.	Observer.
106.28	Spinning disc	Rayleigh, 1882	106.29	Induced discharge	Glazebrook, '88
106.22	" "	Rayleigh and Mrs. Sedgwick, 1883	106.32	Spinning disc	V. Jones, 1894
106.32	Mean result	Rowland, 1887	106.27	" "	Ayrton and V. Jones, 1897
			106.245	" "	Smith, N.P.L., '14

The 1884 "legal" ohm = .9972 intl. ohm; the **B.A. ohm** = .9866 intl. ohm.

THE AMPERE

The electrochemical equivalent of silver is given in milligrams per coulomb (1 ampere for 1 sec.) = 10^{-1} E.M. unit of quantity. **Mean = .00111827 gm./coulomb.** Compare the international ampere (p. 6).

mg. Ag.	Method.	Observer.	mg. Ag.	Method.	Observer.
1.11828	Dynamometer	Kohlrausch, '84	1.11821	Dynamometer	Janet, Laporte, de la Gorce, 1909
1.11827	Current weigher	Corrected 1908 Smith, Mather, and Lowry, 1907	1.11829	"	Do, 1910

E.M.F. OF WESTON CELL

The electromotive force (E) of the Weston (cadmium) cell in volts (10^8 E.M. units) as realized from one of the accepted specifications. The present accepted international value of E is **1.0183 international volts** (see p. 6) at 20° C.

Temperature coefficient.—Over the range 0° to 40°, Wolff (1908) obtained for the E.M.F. at t° —

$$E_t = E_{20} - .0000406(t - 20) - 9.5 \times 10^{-7}(t - 20)^2.$$

E at 20°.	Method.	Observer.	E at 20°.	Method.	Observer.
1.0185	} Intl. ohm and dynamometer	Guthe, 1906	1.01820	Intl. ohm and current weigher	Ayrton, Mather, and Smith, 1908
1.01822		Guillet, 1908	1.01822	" "	Dorsey, 1911
1.01841		Pellat, 1908	1.01834	Intl. ohm and intl. ampere	Jaeger and v. Steinwehr, 1909
1.01869	Intl. ohm and current weigher	Janet, Laporte, Jouaust, 1908			

The E.M.F. of the **Clark cell** = 1.433 volts at 15° C. It diminishes by about 1.2 parts in 1000 for 1° C. rise of temp.

CONVERSION FACTORS

BRITISH INTO METRIC CONVERSION FACTORS

Conversion factors based on the relations given on p. 4. *g* is taken as 981 cm.-sec.⁻². Reciprocals are given for converting metric into British measure.

British.	Metric.	(Reciprocal.)	British.	Metric.	(Reciprocal.)
Length—			Force—		
1 inch	= 2'5400 cm.*	'3937 †	1 poundal	= 13,825 dynes	7'233 × 10 ⁻⁶
1 yard	= '9144 metre*	1'0936	1 pound wgt.	= 4'45 × 10 ⁵ dynes	2'247 × 10 ⁻⁶
1 mile	= 1'6093 km.	'6214	Pressure—		
Area—			1 lb./sq. inch	= 68,971 dynes/cm. ²	1'45 × 10 ⁻⁵
1 sq. inch	= 6'4516 sq. cm	'1550 †	” ”	= 70'31 gm./cm. ²	'01422
Volume—			1 ton/sq. inch	= 1'545 × 10 ⁸ dynes/cm. ²	6'47 × 10 ⁻⁹
1 cubic inch	= 16'387 c.c.	'0610	” ”	= 1'575 k. gm./mm. ²	'6349
1 cubic foot	= 28'317 litre	'03531	Work—		
1 pint	= '5682 litre	1'7598	1 ft.-pound	= 1'356 joules§	'7373
1 gallon	= 4'5460 litre †	'2200 †	Power—		
Mass—			1 horse-power	= '746 k.watt.	1'34
1 grain	= '0648 gram	15'432	Heat—		
1 oz. (avoir.)	= 28'350 grams	'03527	1 B. Th. unit }	= 252'00 calories	'00397
1 lb. ”	= '4536 k. gm.	2'2046	(1 lb., 1° F.)		
1 ton ”	= 1016 k. gm.	'09842	Density—		
Density—			Velocity—		
1 lb./cub. ft.	= '01602 gm./cm. ³	62'43	1 mile/hour	= 44'70 cm./sec.	'02237
Velocity—					

MISCELLANEOUS DATA

CONVENIENT APPROXIMATE RELATIONS		British.	U.States.	
1 yard = 1 metre, less 10%		Std. } yd. at } 62°F. }	{ Stand. } { yd. at } { 59°-6F. }	{ 1 mm. = 10 ⁻³ metre 1 micron, μ = 10 ⁻⁶ " μμ = 10 ⁻⁹ " 1 Å. Ū. = 10 ⁻¹⁰ " 1 mil = 10 ⁻³ inch
2 lbs. = 1 k. gram, "				
2 galls. = 10 litres, "				
1 ton = { 1 tonne				
{ (1000 k. gm.) } less 2%				
SOME BRITISH WEIGHTS AND MEASURES		MATHEMATICAL		
Useful in photography, etc.				
The avoirdupois, troy, and apothecaries grain are the same in weight.				
1 lb. (avoir.) = 7000 grains = 454 grams		π	3'141592654	'49715
1 oz. " = 437½ " = 28'3 "		π ²	9'869604401	'99430
1 oz. (troy) =		1/π	'318309886	1'50285
1 oz. (apothecaries) = 480 " = 31'1 "		√π	1'772453851	'24857
1 fl. drachm = 60 minims = 3'55 c.cs.		1 radian	57°29578	1'75812
1 fl. oz. ⅔ = 8 fl. drachms = 28'41 "		1°	'017453 radian	2'24188
1 pint = 20 fl. ozs. = 568 "		e	2'718281828	'43429
		log. 10	2'302585	'36222
A 10% solution is		To convert		
1 grain in 10 minims of solution		Multiply by		
1 oz. (avoir.) " 10 fl. ozs. "		Common into hyperbolic logs, 2'3026		
2 oz. " " 1 pint "		Hyperbolic " common " '4343		

* Correct to 1 part in a million.

† Correct to 3 parts in a million.

‡ Owing to the definition of the gallon (see p. 4), this number is dependent on assumed buoyancy and temperature corrections.

§ 1 joule = 10⁷ ergs.

|| 1 tonne = 1000 k. g n.

MISCELLANEOUS DATA

MISCELLANEOUS DATA—continued.

BRITISH COINAGE			NAUTICAL		
Coin.	Weight.	Diameter.			
sovereign	8 grams less .15%	2.18 cm.	1 nautical mile	= 6082.66 feet	
penny	$\frac{1}{4}$ oz. (avoir.)	1.2 inch	1 admiralty mile	= 6080 feet	
halfpenny	$\frac{1}{8}$ " "	1.0 " "	1 knot	= 1 nautical mile/hour	
farthing	$\frac{1}{16}$ " "	.8 " "	1 fathom	= 6 feet	
			1 point	= $11\frac{1}{4}^{\circ}$	
10° Centigrade = 50° Fahrenheit, whence the following is convenient for transforming room temperatures:— $5(^{\circ}\text{F.} - 50) = 9(^{\circ}\text{C.} - 10)$				British.	Continental.
			Million . . .	10^6	10^6
			Billion . . .	10^{12}	10^9
			Trillion . . .	10^{18}	10^{12}

VOLUME OF A KILOGRAMME OF PURE WATER

At 4° C. and 760 mm. Values recalculated by Benoît. (*Trav. et Mém. Bur. Intl.*, 14, 1910.) (See p. 4.)

Observer.	C.CS.	Observer.	C.CS.
Lefèvre-Geneau and Fabbroni, 1799 . . .	1000.030	Chaney, 1893	1000.150
Schuckburgh and Kater, 1798 and 1821 . . .	999.525	Guillaume, 1904	1000.039
Svanberg and Berzelius, 1825	999.710	Chappuis, 1907	1000.027
Stampfer, 1831	1000.250	de Lépinau, Benoît, and Buisson, 1907	1000.028
Kupffer, 1842	1000.069		

DENSITIES OF GASES

Supplementary to p. 26. Densities in grams per litre at 0° C., 760 mm., sea-level, and lat. 45°.

Gas.	gms./litre.	Observer.	Gas.	gms./litre.	Observer.
He1782	Watson, <i>J.C.S.</i> , 1910	Ra, Em.	9.727	Gray & Ramsay, <i>P.R.S.</i> 1910
Ne9002	" " "			Baume & Perrot, <i>C.R.</i> , 1909
Kr . . .	3.708	Moore " 1908	CH ₄	.7168	
Xe . . .	5.851	" " "			

C.R., Compt. Rend.; J.C.S., Journ. Chem. Soc.; P.R.S., Proc. Roy. Soc.

PRESSURE COEFFICIENTS OF PV

Pressure coefficient, m , of $p\nu$ for gases at 1 atmosphere and constant temperature; p is the pressure in atmospheres, and ν is the volume. $m = \frac{\partial(p\nu)}{p\nu} \cdot \frac{1}{\partial p}$; m is a measure of the deviation of the gas from Boyle's law.

Air, $m = -.00191$, Regnault.

N, $m = -.000559$

H, $m = +.000772$ } Chappuis, Rayleigh, Leduc, and Sacerdote.

VALUES OF GRAVITY ("g") LONGITUDE AND LATITUDE

Helmert's formula connecting "gravity" with latitude and height is $g = 980.617 - 2.593 \cos 2\lambda - .0003086H$, where λ is the latitude, H is the height in metres above sea-level, and 980.617 cms./sec.² is the value of g attributed to lat. 45° and sea-level. The values of g calculated by this formula are for most places in fair agreement with the observed values. Some discrepancy is found in the vicinity of large mountain ranges, such as the Himalayas.

No absolute standard determination of g has been made in England for many years, but comparisons have been made with Potsdam and Sèvres. For relative measurements, the relation $dg = .0226 dN$ is useful, where N is the number of vibrations which a pendulum makes in a mean solar day of 86,400 mean time seconds. The length (l) of the "seconds" pendulum (*i.e.* 2 secs. period) = $g/\pi^2 = .101321g$. l varies from 99.094 cms. at the equator to 99.620 cms. at the pole.

See Helmert's "Höhere Geodäsie," "Die Grösse der Erde," 1906, and "Die Schwerkraft im Hochgebirge," Clarke's "Geodesy," 1880, Sir Geo. Darwin's "Tides and Kindred Phenomena," Fisher's "Physics of the Earth's Crust," and for recent aspects of the subject, the reports to the triennial International Geodetic Conferences (...1906, 1909...), and the reports of the U.S. Geodetic Survey. (See also p. 13.)

Place.	Longitude E. or W. of Greenwich.	Latitude (λ).	Height (H) above Sea- level.	"g" (calculated).
	° "	° ' "	metres.	cms./sec. ²
Pole	—	90 0 0	—	983.210 *
Equator	—	0 0 0	—	978.024 *
British Isles—				
Aberdeen (Univ.)†.	2 6 38 W	57 8 58 N	21	981.68
Aberystwith	4 4 W	52 25 N	—	981.28 *
Bangor	4 8 W	53 13 N	—	981.35 *
Belfast	5 56 W	54 37 N	—	981.47 *
Birmingham	1 54 W	52 28 N	—	981.28 *
Bristol	2 35 W	51 28 N	—	981.20 *
Cambridge (Univ. Obs.)	0 5 41 E	52 12 52 N	28	981.251
Cardiff	3 10 W	51 28 N	—	981.20 *
Dublin (Trin. Coll.)	6 15 W	53 20 35 N	7 †	981.36
" (R.C.S)	6 40 32 W	53 23 13 N	15	981.36
Dundee (Univ. Coll.)†.	2 58 45 W	56 27 26 N	27 †	981.62
Durham	1 34 56 W	54 46 6 N	—	981.48 *
Edinburgh	3 11 3 W	55 55 28 N	134	981.54
Eskdalemuir (Obs.)	3 12 18 W	55 18 48 N	244	981.45
Glasgow (Univ.)†.	4 17 12 W	55 52 31 N	46	981.56
Greenwich (Obs.)	0 0 0	51 28 38 N	47	981.184
Kew (Obs.)	0 18 46 W	51 28 6 N	5	981.200
Leeds (Univ.)†.	1 33 15 W	53 48 30 N	81	981.38
Liverpool (Univ.)†.	2 57 37 W	53 24 19 N	51	981.35
London (Nat. Phys. Lab.)§.	0 20 11 W	51 25 20 N	10	981.196
" (Univ., S. Kens.)	0 10 23 W	51 29 54 N	14	981.19
" (Univ. Coll.)†.	0 7 57 W	51 31 27 N	28	981.19
Manchester (Univ.)†.	2 14 2 W	53 27 53 N	39	981.37
Newcastle (Armstrong Coll.)	1 36 53 W	54 58 50 N	55	981.48
Nottingham (Univ. Coll.)†.	1 8 45 W	52 57 10 N	58	981.31
Oxford (Radcliffe Obs.)	1 15 39 W	51 45 34 N	65	981.20
Plymouth	4 9 W	50 22 N	—	981.10 *
Portsmouth	1 6 12 W	50 48 3 N	5	981.14
St. Andrews (Univ.)	2 48 W	56 20 N	—	981.62 *
Sheffield (Univ. Obs.)	0 5 50 E	53 23 2 N	—	981.36 *
Stonyhurst (Obs.)	2 28 10 W	53 50 40 N	114	981.37
Africa—				
Bloemfontein	26 40 E	29 0 S	—	979.24 *

* No correction has been applied for height above sea-level.

† Physics laboratory.

§ Teddington.

† Ground floor.

|| Second floor.

GRAVITY

Place.	Longitude E. or W. of Greenwich.	Latitude (λ).	Height (H) above Sea- level.	"g" (calculated).
	° ' "	° ' "	metres.	cms./sec.*
Africa (contd.)—				
Cairo (Observatory)	31 17 14 E	30 4 38 N	33	979'32
Cape Town	18 29 E	33 56 S	12	979'64
Durban	30 40 E	29 40 S	—	979'29*
Johannesburg (Univ. Coll.)	28 7 E	26 11 S	1753	978'49
Mauritius (Roy. Alf. Obs.)	57 33 9 E	20 5 39 S	55	978'63
America—				
Baltimore (Meteorol. Stn.)	76 37 W	39 18 N	23	980'10
Boston (Meteorol. Stn.)	71 4 W	42 21 N	38	980'37
Chicago (Meteorol. Stn.)	87 38 W	41 52 N	251	980'26
Harvard, Camb. (Obs.)	71 7 46 W	42 22 48 N	24	980'37
Jamaica (Montego Bay Obs.)	77 52 22 W	18 24 51 N	69	978'53
Montreal (McGill Obs.)	73 34 39 W	45 30 17 N	57	980'64
New York (Ruthfd. Obs.)	73 59 9 W	40 43 49 N	96	980'20
Philadelphia (Obs.)	75 9 37 W	39 57 8 N	36	980'15
Princeton (N.J.)	74 39 22 W	40 20 58 N	65	980'20
Quebec (Obs.)	71 13 8 W	46 48 21 N	70	980'76
St. Louis (Obs.)	90 12 17 W	38 38 4 N	171	979'99
Toronto (Obs.)	79 23 40 W	43 39 36 N	107	980'46
Washington (Bur. of Stand.)	77 3 59 W	38 56 32 N	102	980'097
Yale, New Haven (Obs.)	72 55 8 W	41 19 22 N	32	980'28
Asia—				
Bombay (Obs.)	72 48 56 E	18 53 45 N	10	978'57
Calcutta (Surv. Office)	88 21 30 E	22 32 54 N	6	978'76
Hong Kong (Obs.)	114 10 28 E	22 18 13 N	33	978'76
Madras (Obs.)	80 14 54 E	13 4 8 N	7	978'29
Australasia—				
Adelaide (Obs.)	138 35 8 E	34 55 39 S	430	979'68
Brisbane (Obs.)	153 1 36 E	27 28 S	42	979'12
Melbourne (Obs.)	144 58 32 E	37 49 53 S	28	979'97
Perth.	115 52 E	31 57 S	14	979'47
Sydney (Obs.)	151 12 23 E	33 51 41 S	44	979'63
Wellington (Obs.), N.Z.	174 46 37 E	41 18 1 S	43	980'27
Europe—				
Berlin (Reichsanstalt) †	13 19 E	52 31 N	30	981'287
Christiania (Obs.)	10 43 23 E	59 54 44 N	25	981'90
Copenhagen (Obs.)	12 31 40 E	55 41 13 N	14	981'56
Geneva (Obs.)	6 9 11 E	46 11 59 N	374	980'61
Leyden (Obs.)	4 29 3 E	52 9 20 N	6	981'26
Paris (Obs.)	2 20 14 E	48 50 11 N	59	980'95
" (Bureau Intl.) †	2 13 10 E	48 49 53 N	70	980'951
Potsdam (Astron. Inst.)	13 3 59 E	52 22 56 N	94	981'249
Rome (Coll. Obs.)	12 28 53 E	41 53 54 N	59	980'32
St. Petersburg (Acad. Obs.)	30 18 22 E	59 56 30 N	3	981'91
Vienna (Impl. Obs.)	16 20 21 E	48 12 47 N	—	980'91*
Zurich (Poly. Obs.)	8 33 4 E	47 22 40 N	468	980'69

* No correction applied for height above sea-level.

† Charlottenburg.

‡ Sèvres.

DISTANCES ON THE EARTH'S SURFACE
(See Ball's "Spherical Astronomy," 1909.)

At Lat.	Miles per degree of		At Lat.	Miles per degree of		At Lat.	Miles per degree of	
	Longitude.	Latitude.		Longitude.	Latitude.		Longitude.	Latitude.
0	69'15	68'69	40	53'05	69'00	60	34'66	69'21
10	68'11	68'70	45	48'99	69'05	70	23'73	69'32
20	65'01	68'77	50	44'54	69'10	80	12'05	69'38
30	59'94	68'88	55	39'75	69'16	90	0	69'39

SIZE AND SHAPE OF THE EARTH

The spheroid of revolution which most nearly approximates to the earth, has the following dimensions :—
[1 kilom. = .6214 mile.]

Observer.	Equatorial radius, a .	Polar radius, b .	Ellipticity, $(a-b)/a$.
Bessel, 1841 . . .	6,377,397 metres	6,356,079 metres	$1/299.2$
Clarke, 1866 . . .	8,206 "	584 "	$1/295.0$
" 1880 . . .	8,249 "	515 "	$1/293.5$
Helmert, 1906 * .	8,200 "	818 "	$1/298.3$
U.S. Survey, 1906 †	8,388 " ‡	909 "	$1/297.0$

* "Die Grosse der Erde."

† "The Figure of the Earth," 1909, and Supplement, 1910; U.S. Coast and Geodetic Survey.

‡ 3963.339 miles. || 3949.992 miles.

MEAN DENSITY OF THE EARTH

(See Poynting's "Mean Density of the Earth," 1893.)

Observer.	Density.
Common Balance Method.	
Poynting, 1878	5.493
Richarz and Krigar-Menzel, 1898	5.505
Torsion Balance Method.	
Cavendish, 1798	5.45
Boys, <i>Phil. Trans.</i> , 1895 . .	5.527
Braun, 1896	5.527
Eötvös, 1896	5.534
Mean density of surface . . .	2.65

SUN

The mean equatorial solar parallax (Hinks, 1909) } = $8''.807$

Whence mean distance } = $\begin{cases} 1.494 \times 10^{11} \\ \text{metres} \\ 9.282 \times 10^7 \\ \text{miles} \end{cases}$

Mean time taken by light to travel from sun to earth } = 498.2 secs.

MOON

Mean distance from earth to moon } = $\begin{cases} 60.27 \times \text{earth's} \\ \text{radius} \end{cases}$
 Mass of the moon } = $\begin{cases} (1/81.53) \times \\ \text{earth's mass} \end{cases}$ (Hinks, 1909)
 Inclination of moon's orbit to ecliptic } = $5^\circ 8' 43''$

Mean polar quadrant } = $10,002,100$ metres *
 Volume of earth = 1.083×10^{21} metres³ *
 Mass of earth = 5.98×10^{27} grams †
 = 5.87×10^{21} tons
 Area of land = 1.45×10^{18} cm.²
 Area of ocean = 3.67×10^{18} cm.²
 Mean depth of ocean (Murray) } = 3.85×10^6 cm.
 Volume of ocean = 1.41×10^{24} cm.³
 Mass of ocean = 1.45×10^{24} grms.

Constant of Gravitation (G in law of attraction) = 6.658×10^{-8} c.g.s.

Obliquity of the Ecliptic to the equator = $23^\circ 27' 4''.04$ in 1909, subject to a small fluctuation by nutation, and a slow continuous decline of $46''.84$ per century.

Constant of aberration of a star is theoretically equal to (Earth's orbital velocity)/(velocity of light) = $20''.43 \pm''.03$ (Renan and Ebert, 1905).

Constant of precession, i.e. annual precessional increase of the longitude of a star = $50''.2564 +''.0002225t$, where t is the interval in years from 1900 (Newcomb).

* Mean of Helmert and U.S. Survey.

† Using Boys' and Braun's result for density.

SOLAR SYSTEM

ELEMENTS OF THE SOLAR SYSTEM

8''·806 is taken as the equatorial horizontal solar parallax from the observations of the asteroid Eros in 1920-1; 5'527 is adopted as the Earth's mean density (Boys, 1895; Braun, 1896). The constants for Mercury are those adopted by Stroobant and Backlund (1909). The value of the mass of Jupiter is that obtained by Cookson (1908). The time of rotation of Venus is that suggested by Hansky and Stefánik (1907). (See Newcomb's "Spherical Astronomy" and Ball's "Spherical Astronomy.")

Name.	Equatorial Semi-diameter.			Mass Earth = 1	Mean Density.		Gravity at Surf. Earth = 1	No. of Satellites.†
	Angular.*	Miles.	Earth = 1		Earth = 1	Water = 1		
Sun . .	16' 11.8"	432,890	109.2	329,390	.25	1.39	27.61	—
Mercury	3'08"	1387	.350	.34	.88	4.86	.28	0
Venus .	8'40"	3783	.955	>.818	>.94	>5.20	>.91	0
Earth .	8'80"	3963.3	1.000	1.000	1.00	5.527	1.00	1 (D)
Mars . .	4'68"	2108	.532	.106	.071	3.90	.38	2 (D)
Jupiter .	1 37'36"	43850	11.06	314.50	.25	1.36	2.57	8 (7 D; 1 R)
Saturn .	1 24'75"	38170	9.63	94.07	.12	.63	1.01	10 (9 D; 1 R)
Uranus .	34'28"	15440	3.90	14.40	.24	1.34	.95	4 (R)
Neptune	36'56"	16470	4.15	16.72	.23	1.28	.97	1 (R)

Name.	Inclination of Equator to Orbit.	Time of Axial Rotation.	Semi-major Axis of Orbit.		Sidereal Period.		
			Earth = 1.		Millions of Miles.	Mean Solar Days.	Julian Years.
Sun . .	0' 15"	d h m 25 9 7	—	—	—	—	—
Mercury.	?	?	3870986	Bode's Law 4 = (0+4)	36.0	87.9693	.24
Venus .	?	23 40 (?)	7233315	7 = (3+4)	67.2	224.7008	.62
Earth .	23 27 8	23 56 4.09	1.0000000	10 = (6+4)	92.9	365.2564	1.00
Mars . .	24 52	24 37 22.74	1.523688	16 = (12+4)	141.6	686.9797	1.88
Asteroids	—	—	2.55 to 2.85	28 = (24+4)	237 to 265	—	—
Jupiter .	3 5	9 56 ±	5.202803	52 = (48+4)	483.3	4332.588	11.86
Saturn .	26 49	10 15 ±	9.538844	100 = (96+4)	886.2	10759.20	29.46
Uranus .	?	13 ?	19.19098	196 = (192+4)	1782.8	30586.29	83.74
Neptune	27 ?	?	30.07067	—	2793.5	60187.65	164.78

Name.	Ellipticity of Planet. §	Mean Daily Motion in Orbit.	Longitude of Perihelion.	Longitude of Ascending Node. ¶	Inclination of Orbit to Ecliptic.	Eccentricity of Orbit. **
Mercury.	?	4 5 32.4	75 53 59	47 8 45	7 0 10	.205614
Venus .	?	1 36 7.7	130 9 50	75 46 47	3 23 37	.006821
Earth .	1/298.3	59 8.2	101 13 15	0 0 0	0 0 0	.016751
Mars . .	1/270 ?	31 26.5	334 13 7	48 47 9	1 51 1	.093309
Jupiter .	1/17	4 59.1	12 36 20	99 26 42	1 18 42	.048254
Saturn .	1/9	2 0.5	90 48 32	112 47 12	2 29 39	.056061
Uranus .	1/95.2	42.2	169 2 56	73 29 25	0 46 22	.047044
Neptune	?	21.5	43 45 20	130 40 44	1 46 45	.008533

* This is the angle subtended by the semi-diameter at a distance equal to the Earth's mean distance from the Sun.

† The inclination of the plane of the Sun's equator to the plane of the ecliptic.

‡ D means direct; R, retrograde.

§ The ellipticity = $(a-b)/a$, where a is the major axis and b the minor axis of the spheroid of revolution. The value given for the Earth is Helmert's (p. 13).

|| Perihelion is the point in the orbit nearest the Sun. Longitude is the angular distance from the first point of Aries (see p. 3), measured along the ecliptic.

¶ A node is one of the two points at which a planet's orbit intersects the plane of the ecliptic. At the ascending node the planet passes from south to north of the ecliptic.

** The eccentricity = $\sqrt{(a^2 - b^2)}/a$, where a and b are the major and minor axes of the orbit.

EQUATION OF TIME

(+) means that the equation of time has to be added to the apparent solar time (*i.e.* sundial time) to give the mean solar or clock time (see p. 3). (M) = maximum or minimum. The values below vary by a few seconds from year to year. $C = D + E$, where C = clock time, D = dial time, and E = equation of time.

Date.	Equation of time.	Date.	Equation of time.	Date.	Equation of time.	Date.	Equation of time.
	m. s.		m. s.		m. s.		m. s.
Jan. 1	+ 3 11	April 1	+ 4 1	July 1	+ 3 32	Oct. 16	- 14 20
" 16	+ 9 33	" 16	0 0	" 26	+ 6 18 (M)	Nov. 3	- 16 21 (M)
Feb. 1	+ 13 37	May 1	- 2 57	Aug. 16	+ 4 11	" 16	- 15 10
" 12	+ 14 25 (M)	" 14	- 3 49 (M)	Sept. 1	0 0	Dec. 1	- 10 56
Mar. 1	+ 12 34	June 1	- 2 27	" 16	- 5 6	" 12	- 6 15
" 16	+ 8 51	" 15	0 0	Oct. 1	- 10 16	" 25	0 0

PARALLAXES OF STARS

The **proper motion** of a star is its real change of place arising from the actual motion of the star itself.

The **annual parallax** is the angle between the direction in which a star appears as seen from the earth and the direction in which it would appear if it could be observed from the centre of the sun.

A **light-year** is the distance that light travels in one year (see p. 69).

Star and Magnitude.	Proper motion per year.	Annual parallax.	Distance.	
			Sun's dist. = 1	Light-years.
α Centauri (2)	3.7	$.75 \pm .01$	$.28 \times 10^6$	4.4
21185 Lalande (7.5)	7.3	$.48 \pm .02$.43	6.8
61 Cygni (4.8)	5.2	$.37 \pm .02$.56 "	8.8
Sirius (-1.4)	1.3	$.37 \pm .01$.56 "	8.8
Procyon (.5)	1.3	.31	.69 "	11
Altair (.9)7	$.28 \pm .02$.74 "	12
Aldebaran (1.1)2	$.17 \pm .02$	1.4 "	22
Capella (.2)4	$.12 \pm .02$	1.7 "	27
Vega (.1)4	$.12 \pm .02$	1.7 "	27
1830 Groombridge (6.4)	7.0	$.10 \pm .02$	2.0 "	33
Polaris (2.1)	0.0	$.07 \pm .02$	3.0 "	47
Arcturus (.2)	2.3	.024	8.7 "	140

SYSTEMATIC MOTIONS OF THE STARS

The apparent proper motions of the stars show drifts in two directions. The assigned positions of the apices of these directions are:—

Computer.	Stream I.		Stream II.	
	R.A.	Dec.	R.A.	Dec.
Kapteyn, 1904	85°	- 11°	260°	- 48°
Eddington	90°	- 19°	292°	- 58°
Dyson	94°	- 7°	240°	- 74°

STANDARD TIMES

Referred to Greenwich time.

Gt. Britain, France, Portugal, Belgium, Spain	Greenwich time
Ireland	"
Austria, Denmark, Germany, Italy, Norway, Switzerland	1 hour fast
British South Africa, Egypt, Turkey	1½ or 2 hours fast
Japan	9 hours fast
Australia	8, 9, or 10 hours fast
New Zealand	11½ " "
Canada and United States	4, 5, 6, 7, or 8 hours slow

SCREWS

SCREWS

It is customary for British metal screws, of $\frac{1}{4}$ -inch diameter and above, to have a Whitworth thread, for smaller sizes a British Association thread. In the Whitworth thread the angle between the slopes is 55° , in the B.A. thread 47.5° .

The **pitch** is the distance between adjoining crests (say) of the same thread measured parallel to the axis of the screw. It is the reciprocal of the number of turns per inch or mm. as the case may be. The **full diameter** is the maximum over-all diameter.

Micrometer screws are made with some multiple or sub-multiple of 100 threads to the inch or mm.

"**Wood screws**" of iron or brass are numbered as follows: No. 0 has a diameter of .05 inch, each succeeding number adding .014 inch to the diameter of the screw: this applies to all lengths. The length of countersunk screws is measured over all; that of round-headed screws, from under the head. [1 inch = 25.4 mm.]

STANDARD WHITWORTH.				BRITISH ASSOCIATION.								
Full di- ameter	Threads to inch.	Full di- ameter.	Threads to inch.	No.	Full di- ameter.	Pitch.	No.	Full di- ameter.	Pitch.	No.	Full di- ameter	Pitch.
inch.		inch.			mm	mm.		mm.	mm.		mm.	mm.
1 $\frac{3}{8}$	5	$\frac{3}{4}$	10	0	6.0	1.0	9	1.9	.39	18	.62	.15
1 $\frac{1}{2}$	5	$\frac{11}{16}$	11	1	5.3	.9	10	1.7	.35	19	.54	.14
1 $\frac{3}{4}$	6	$\frac{9}{16}$	11	2	4.7	.81	11	1.5	.31	20	.48	.12
1 $\frac{7}{8}$	6	$\frac{13}{16}$	12	3	4.1	.73	12	1.3	.28	21	.42	.11
1 $\frac{1}{4}$	7	$\frac{1}{2}$	12	4	3.6	.66	13	1.2	.25	22	.37	.10
1 $\frac{1}{8}$	7	$\frac{7}{8}$	14	5	3.2	.59	14	1.0	.23	23	.33	.09
1	8	$\frac{5}{8}$	16	6	2.8	.53	15	.9	.21	24	.29	.08
$\frac{7}{8}$	9	$\frac{15}{16}$	18	7	2.5	.48	16	.79	.19	25	.25	.07
$\frac{1}{2}$	10	$\frac{1}{4}$	20	8	2.2	.43	17	.70	.17			

MOMENTS OF INERTIA

M = mass of body. (See A. M. Worthington, "Dynamics of Rotation." London.)

Body.	Axis of rotation.	Moment of inertia.
Uniform thin rod (length l)	(1) Through centre, perpendicular to length	$M \frac{l^2}{12}$
	(2) Through end, perpendicular to length	$M \frac{l^2}{3}$
Rectangular lamina (sides a and b)	(1) Through centre of gravity, perpendicular to plane	$M \frac{a^2 + b^2}{12}$
	(2) Through centre of gravity, parallel to side b	$M \frac{a^2}{12}$
Circular lamina (radius r)	(1) Through centre, perpendicular to plane	$M \frac{r^2}{2}$
	(2) Any diameter	$M \frac{r^2}{4}$
Solid cylinder (radius r ; length l)	(1) Axis of cylinder	$M \frac{r^2}{2}$
	(2) Through centre of gravity, perpendicular to axis of cylinder	$M \left(\frac{l^2}{12} + \frac{r^2}{4} \right)$
Hollow cylinder (external and internal radii R and r ; length l)	(1) Axis of cylinder	$M \frac{R^2 + r^2}{2}$
	(2) Through centre of gravity, perpendicular to axis	$M \left(\frac{l^2}{12} + \frac{R^2 + r^2}{4} \right)$
Solid sphere (radius r)	Through centre	$M \frac{2r^2}{5}$
Hollow sphere (external and internal radii R and r)	Through centre	$M \left(\frac{2}{5} \frac{R^5 - r^5}{R^3 - r^3} \right)$
Anchor ring (mean radius of ring R ; radius of cross-section r)	(1) Through centre, perpendicular to plane of ring	$M \left(R^2 + \frac{3r^2}{4} \right)$
	(2) Any diameter	$M \left(\frac{R^2}{2} + \frac{5r^2}{8} \right)$

VOLUME CALIBRATION OF VESSELS BY WATER OR MERCURY

Volume content of vessel at $t^\circ \text{C.} = V_t = W_t v_t \equiv w_t(f)$, where—

w_t = observed weight in grams (against brass weights in air) of contained water (or mercury) at $t^\circ \text{C.}$

W_t = weight of such liquid *in vacuo* (i.e. corrected for buoyancy in air).

v_t = volume of 1 gram of liquid at $t^\circ \text{C.}$

(f) is a factor which introduces the buoyancy and specific volume corrections.

The following table of values of the factor (f) is based on tables on pp. 19 and 22.

Temp. (t) of weighing	10° C.	11°	12°	13°	14°	15°	16°	17°
Value of $\left\{ \begin{array}{l} \text{H}_2\text{O} \\ \text{factor } (f) \end{array} \right\} \text{Hg}$	1'00133 '073683	1'00143 '073697	1'00154 '073710	1'00166 '073724	1'00179 '073737	1'00193 '073750	1'00209 '073764	1'00226 '073777
Temp. (t) of weighing	18°	19°	20°	21°	22°	23°	24°	25°
Value of $\left\{ \begin{array}{l} \text{H}_2\text{O} \\ \text{factor } (f) \end{array} \right\} \text{Hg}$	1'00244 '073790	1'00263 '073804	1'00283 '073817	1'00305 '073831	1'00327 '073844	1'00350 '073857	1'00375 '073871	1'00400 '073884

The above gives the volume content V_t of the vessel at the temperature of weighing, $t^\circ \text{C.}$ At any other temperature, t' , the volume $V_{t'} = V_t \{1 + \gamma(t' - t)\} \equiv V_t(F)$, where γ is the coefficient of cubical expansion of the material of the vessel. Values of the factor (F) for **glass vessels** ($\gamma = .00025$) are tabulated below.

($t' - t$)	2° C.	4°	6°	8°	-2° C.	-4°	-6°	-8°
Value of factor (F)	1'00005	1'00010	1'00015	1'00020	'99995	'99990	'99985	'99980

Example.—Weight of water contained in a vessel at $10^\circ \text{C.} = 10$ grams: thence volume of vessel at $10^\circ \text{C.} = 10 \times 1'00133$ c.c.s. The same vessel, if of glass, would contain at $16^\circ \text{C.}, 10 \times 1'00133 \times 1'00015 = 10'0148$ c.c.s.

CAPILLARITY CORRECTIONS OF MERCURY COLUMNS

The height of the meniscus and the value of the capillary depression depend on the bore of the tubing, on the cleanliness of the mercury, and on the state of the walls of the tube. The correction is negligible for tubes with diameters greater than about 25 mms. The table below gives the amount of the correction (which has to be added to the height) for various diameters of glass tubing and meniscus heights. (Mendeléeff and Gutkowsky, 1877. See also Scheel and Heuse, *Ann. d. Phys.*, 33, 1910.)

Bore of tube.	Height of meniscus in mms.								Bore of tube.	Height of meniscus in mms.					
	.4	.6	.8	1.0	1.2	1.4	1.6	1.8		.8	1.0	1.2	1.4	1.6	1.8
mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.
4	.83	1'22	1'54	1'98	2'37	—	—	—	9	.21	.28	.33	.40	.46	.52
5	.47	.65	.86	1'19	1'45	1'80	—	—	10	.15	.20	.25	.29	.33	.37
6	.27	.41	.56	.78	.98	1'21	1'43	—	11	.10	.14	.18	.21	.24	.27
7	.18	.28	.40	.53	.67	.82	.97	1'13	12	.07	.10	.13	.15	.18	.19
8	—	.20	.29	.38	.46	.56	.65	.77	13	.04	.07	.10	.12	.13	.14

BAROMETRY

REDUCTION OF BAROMETER READINGS TO 0° C.

Corrected height $H_0 = H \left\{ 1 - \frac{(\beta - \alpha)l}{(1 + \beta l)} \right\}$, where H and l are the observed height and temperature of the barometer, $\beta = .0001818$ (Regnault), the coefficient of cubical expansion of mercury; $\alpha = .0000085$, the coefficient of linear expansion of glass, or .000184 for brass. Hydrogen temperature scale. (After Broch, Inter. Bur. Weights and Measures.)

(In standard English barometry the mercury is reduced to 32° F., and the scale to 62° F. In the table below, both are reduced to the ice point.)

Temp. (°).	Correction in mms. to be subtracted.									
	GLASS SCALE.					BRASS SCALE.				
	Uncorrected height in mms.					Uncorrected height in mms.				
	700	720	740	760	780	700	720	740	760	780
	mm.					mm.				
2° C.	.24	.25	.26	.26	.27	.23	.24	.24	.25	.25
4	.43	.49	.51	.53	.54	.46	.47	.48	.50	.51
6	.73	.75	.77	.79	.81	.69	.71	.72	.74	.76
8	.97	.99	1.02	1.05	1.08	.91	.94	.97	.99	1.02
10	1.21	1.25	1.28	1.31	1.35	1.14	1.17	1.21	1.24	1.27
12	1.45	1.49	1.53	1.58	1.62	1.37	1.41	1.45	1.49	1.53
14	1.69	1.74	1.79	1.84	1.89	1.60	1.64	1.69	1.73	1.78
16	1.94	1.99	2.05	2.10	2.16	1.82	1.88	1.93	1.98	2.03
18	2.18	2.24	2.30	2.36	2.43	2.05	2.11	2.17	2.23	2.29
20	2.42	2.49	2.56	2.62	2.69	2.28	2.34	2.41	2.47	2.54
22	2.66	2.73	2.81	2.89	2.96	2.51	2.58	2.65	2.72	2.79
24	2.90	2.98	3.06	3.15	3.23	2.73	2.81	2.89	2.97	3.05
26	3.14	3.23	3.32	3.41	3.50	2.96	3.04	3.13	3.21	3.30
28	3.38	3.47	3.57	3.67	3.77	3.19	3.28	3.37	3.46	3.55
30	3.62	3.72	3.83	3.93	4.03	3.41	3.51	3.61	3.71	3.80
32	3.86	3.97	4.08	4.19	4.30	3.64	3.74	3.85	3.95	4.05
34	4.10	4.21	4.33	4.45	4.57	3.87	3.98	4.09	4.20	4.31

REDUCTION OF BAROMETER READINGS TO LAT. 45° AND SEA-LEVEL

It is a convention to take " g " at lat. 45° and sea-level as the standard value for "gravity." The corrections below result from the variation of " g " with latitude and height above sea-level (see p. 11). The barometer correction for $\text{latitude} = \frac{H_0}{760}(\text{C})$, has to be subtracted from the temperature-corrected barometer reading H_0 for latitudes between 0° and 45°; and added for latitudes from 45° to 90°.

Latitude	0° 90°	5° 85°	10° 80°	15° 75°	20° 70°	25° 65°	30° 60°	35° 55°	40° 50°	45° 45°
C	mm. 1.97	1.94	1.85	1.70	1.51	1.27	.98	.67	.34*	.00

The correction of the barometer due to diminution of gravity with increasing height above sea-level amounts to about .24 mm. of mercury per 1000 metres above sea-level. The correction has to be subtracted from the observed reading.

* London, 45.

REDUCTION OF WEIGHINGS TO VACUO

The buoyancy correction = $M\sigma(1/\Delta - 1/\rho) = M\hat{k}$, where M is the apparent mass in grams of the body in air, σ is the density of air (= '0012) in grams per c.c., Δ is the density of the body, ρ is the density of the weights. The correction is true to 4% for the following limits: 740 mm. press., 1° to 22°; 760 mm., 8° to 29°; 780 mm., 15° to 35°. If the correction is required more accurately, multiply the value of \hat{k} given below by σ'/σ , where σ' is the true density of the air for the temp. and press. at the time of the weighing (for σ' , see p. 25). The corrections for quartz weights are the same as for Al. + means corⁿ. to be added to observed weight.

Density of Body weighed Δ .	Correction Factor (\hat{k}) in Milligrams.			Density of Body weighed Δ .	Correction Factor (\hat{k}) in Milligrams.		
	Brass wghts. $\rho = 8.4$.	Pt wghts. $\rho = 21.5$.	Al wghts. $\rho = 2.65$.		Brass wghts. $\rho = 8.4$.	Pt wghts. $\rho = 21.5$.	Al wghts. $\rho = 2.65$.
.5	+ 2'26	+ 2'34	+ 1'95	1'6	+ '61	+ '69	+ '30
.55	+ 2'04	+ 2'13	+ 1'73	1'7	+ '56	+ '65	+ '25
.6	+ 1'86	+ 1'94	+ 1'55	1'8	+ '52	+ '62	+ '21
.65	+ 1'70	+ 1'79	+ 1'39	1'9	+ '49	+ '58	+ '18
.7	+ 1'57	+ 1'66	+ 1'26	2	+ '46	+ '54	+ '15
.75	+ 1'46	+ 1'55	+ 1'15	2'5	+ '34	+ '43	+ '03
.8	+ 1'36	+ 1'44	+ 1'05	3	+ '26	+ '34	+ '05
.85	+ 1'27	+ 1'36	+ '96	3'5	+ '20	+ '29	+ '11
.9	+ 1'19	+ 1'28	+ '88	4	+ '16	+ '24	+ '15
.95	+ 1'12	+ 1'21	+ '81	5	+ '10	+ '19	+ '21
1	+ 1'06	+ 1'14	+ '75	6	+ '06	+ '14	+ '25
1'1	+ '95	+ 1'04	+ '64	8	+ '01	+ '09	+ '30
1'2	+ '86	+ '94	+ '55	10	+ '02	+ '06	+ '33
1'3	+ '78	+ '87	+ '47	15	+ '06	+ '03	+ '37
1'4	+ '71	+ '80	+ '40	20	+ '08	+ '004	+ '39
1'5	+ '66	+ '75	+ '35	22	+ '09	+ '001	+ '40

REDUCTION OF GASEOUS VOLUMES TO 0° AND 760 MMS. PRESSURE

Corrected volume $v_0 = \{v/(1 + '00367t)\} \cdot p/760$, where v , t , and p are the observed volume, temp., and pressure (in mms. of mercury) of the gas respectively. $g = 980.62$ cms. per sec². The coefficient '00367 observed by Regnault.

Values of $(1 + '00367t)$.

Temp. (t).	0	1	2	3	4	5	6	7	8	9
0° C.	1'0000	1'0037	1'0073	1'0110	1'0147	1'0183	1'0220	1'0257	1'0294	1'0330
10	0367	0404	0440	0477	0514	0550	0587	0624	0661	0697
20	0734	0771	0807	0844	0881	0917	0954	0991	1028	1064
30	1101	1138	1174	1211	1248	1284	1321	1358	1395	1431
40	1468	1505	1541	1578	1615	1651	1688	1725	1762	1798
50	1835	1872	1908	1945	1982	2018	2055	2092	2129	2165
60	2202	2239	2275	2312	2349	2385	2422	2459	2496	2532
70	2569	2606	2642	2679	2716	2752	2789	2826	2863	2899
80	2936	2973	3009	3046	3083	3119	3156	3193	3230	3266
90	3303	3340	3376	3413	3450	3486	3523	3560	3597	3633
100	3670	3707	3743	3780	3817	3853	3890	3927	3964	4000
110	4037	4074	4110	4147	4184	4220	4257	4294	4331	4367

Values of $p/760$

Press. (p).	0	1	2	3	4	5	6	7	8	9
700 mm.	'9211	'9224	'9227	'9250	'9263	'9276	'9289	'9303	'9316	'9329
710	'9342	'9355	'9368	'9382	'9395	'9408	'9421	'9434	'9447	'9461
720	'9474	'9487	'9500	'9513	'9526	'9539	'9553	'9566	'9579	'9592
730	'9605	'9618	'9632	'9645	'9658	'9671	'9684	'9697	'9711	'9724
740	'9737	'9750	'9763	'9776	'9789	'9803	'9816	'9829	'9842	'9855
750	'9868	'9882	'9895	'9908	'9921	'9934	'9947	'9961	'9974	'9987
760	1'0000	1'0013	1'0026	1'0039	1'0053	1'0066	1'0079	1'0092	1'0105	1'0118
770	1'0132	1'0145	1'0158	1'0171	1'0184	1'0197	1'0211	1'0224	1'0237	1'0250

DENSITIES

DENSITIES OF THE ELEMENTS

Average densities of liquid and solid elements in grams per c.c. at ordinary temperature unless otherwise stated. For gaseous densities see p. 26. The density of a specimen may depend considerably on its state and previous treatment, *e.g.* the density of a cast metal is increased by drawing, rolling, or hammering.

Element.	Density.	Element.	Density.	Element.	Density.
Aluminium . . .	2.70	Indium	7.12	Samarium	7.8
Antimony . . .	6.62	Iodine	4.95	Scandium	(?)
Argon (liq.) . .	1.4/-185°	Iridium	22.41	Selenium, amorph.	4.8
Arsenic	5.73	Iron (pure) . . .	7.86	" cryst. . .	4.5
Barium	3.75	Krypton (liq.) . .	2.16	" liq. . . .	4.27
Beryllium . . .	1.93	Lanthanum	6.12	Silicon	c. 2.3
Bismuth	9.80	Lead	11.37	Silver	10.5
Boron	2.5 (?)	Lithium	5.34	Sodium971
Bromine	3.102/25°	Magnesium	1.74	Strontium	2.54
Cadmium	8.64	Manganese	7.39	Sulphur, rhombic	2.07
Cæsium	1.87	Mercury (see p. 22)	13.56/15°	" monoclinic	1.96
Calcium	1.55/29°	Molybdenum . . .	10.0	" amorphous	1.92
Carbon—		Neodymium	6.96	" liquid 113°	1.81
Diamond . . .	3.52	Neon (liq.)	(?)	Tantalum	16.6
Graphite . . .	2.3	Nickel	8.9	Tellurium	6.25
Cerium	6.92	Niobium	12.75	Terbium	(?)
Chlorine (liq.) .	2.49/0°	Nitrogen (liq.) . .	7.79/-196°	Thallium	11.9
Chromium	6.50	Osmium	22.5	Thorium	11.3
Cobalt	8.6	Oxygen (liq.) . . .	1.27/-235°	Tin	7.29
Copper	8.93	Palladium	11.4	Titanium	3.54
Erbium	4.77 (?)	Phosphorus, red .	2.20	Tungsten	18.8
Fluorine (liq.) .	1.11/-187°	" yellow . . .	1.83	Uranium	18.7
Gadolinium . . .	(?)	Platinum	21.50	Vanadium	5.5
Gallium	5.95	Potassium	8.62	Xenon (liq.) . . .	3.5
Germanium . . .	5.47	Praseodymium . .	6.48	Ytterbium	(?)
Gold	19.32	Radium	(?)	Yttrium	3.8(?)
Helium (liq.) . .	.12/B.P.	Rhodium	12.44	Zinc	7.1
Hydrogen (liq.) .	.07/B.P.	Rubidium	1.532	Zirconium	4.15
" "086/M.P.	Ruthenium	12.3		

The densities of the alkali metals Li, Na, K, Rb, Cs are due to Richards and Brink, 1907; of He at -268° 6, Onnes, 1908; of W, Gin, 1908; of Ta, Nb, and Th, von Bolton, 1905, 1907, 1908; of Ca, Goodwin, 1904; of Rh and Ir, Holborn, Henning, and Austin, 1904; of Br, Andrews and Carlton, 1907.

DENSITIES OF COMMON SUBSTANCES

Average densities in grams per c.c. at ordinary temperatures. For densities of acids, alkalis, and other solutions, see pp. 23 *et seq.*; of "chemical compounds," p. 109; of gases, p. 26; of other minerals, p. 126.

Substance.	Density.	Substance.	Density.	Substance.	Density.
Metals & Alloys.		Coins (English)		Woods (seasoned).	
Iron, cast	7.1-7.7	" silver \$. . .	10.31	Ash; mahogany . .	6-8
" wrought . .	7.8-7.9	Constantan (Eu- } . . .	8.88	Bamboo	c. 4
" wire	7.7	reka) 		Beach; oak; teak .	7-9
Steel	7.7-7.9	German silver ¶ . .	8.5-8.9	Box	9-11
Brass (ordy.) * .	8.4-8.7	Gunmetal	8.0-8.4	Cedar	5-6
Brass weights . .	c. 8.4	Magnalium ** . . .	c. 2	Ebony	1.1-1.3
Bronze (Cu, Sn)	8.7-8.9	Manganin †† . . .	8.5	Lignum vitæ . . .	1.2-1.3
Coins (English)		Phosphor bronze ††	8.7-8.9	Pitchpine; walnut	6-7
" bronze † . .	8.96	Platinoid §§ . . .	c. 9	Red pine (deal) .	5-7
" gold ‡ . . .	17.72	Pt (90), Ir (10) .	21.62	White pine	4-5

* c. 66 Cu, 34 Zn. † 95 Cu, 4 Sn, 1 Zn. ‡ 91½ Au, 8½ Cu. § 92½ Ag, 7½ Cu. || 60 Cu, 40 Ni.
¶ 60 Cu, 15 Ni, 25 Zn. ** c. 70 Al, 30 Mg. †† 84 Cu, 12 Mn, 4 Ni. ‡‡ 92½ Cu, 7 Sn, ½ P.
§§ Described as German silver with a little tungsten.

DENSITIES OF COMMON SUBSTANCES (contd.)

Substance.	Density.	Substance.	Density.	Substance.	Density.
Minerals, etc.		Liquids.		Gelatine	1.27
Agate; slate . . .	2.5-2.7	Glycerine	1.26	Glass, flint	2.9-4.5
Asbestos	3.0	Methylated spirit83	" crown ;	2.4-2.6
" board	1.2	Milk	c. 1.03	" window }	
Carbon (see above)		Naphtha85	" Jena	(see p. 74.)
Charcoal3-6	Oil, castor97	Ice (Roth, 1908), 0°	.9168
Coal	1.2-1.5	" linseed91-.93	" (Vincent, '02), 0°	.9160
" anthracite . . .	1.4-1.8	" lubricating90-.92	Indiarubber (pure)	.91-.93
Coke	1.0-1.7	" olive ; palm91-.93	Ivory	1.8-1.9
Gas carbon	1.9	" paraffin	c. .8	Leather85-1
Emery	4.0	Petrol68-.72	Paper7-1.1
Granite	2.5-3	Sea-water	1.01-1.05	Pitch	c. 1.1
Marble	2.5-2.8	Turpentine87	Porcelain	2.2-2.4
Masonry	c. 2	Vinegar	1.02	Resin	c. 1.1
Pumice (natural) . .	.4-9	Miscellaneous.		Red fibre	1.45
Quartz	2.66	Amber	1.1	Snow (loose) . . .	c. .12
Silica, fused		Bone	1.8-2.0	Tar	1.02
" transparent . . .	2.21	Butter, lard92-.94	Wax, soft paraffin .	.87-.88
" translucent . . .	2.07	Celluloid	1.4	" hard88-.93
Sand (silver)	2.63	Cork22-.26	" white ; bees-	.95-.96
Sandstone ; kaolin	2.2-2.3	Ebonite	1.3	" sealing	c. 1.8
				" soft red	c. 1.0

DENSITY DETERMINATION CORRECTIONS

In the determination of the density of a body by weighing in water, the true density (corrected for air buoyancy and water density) is given by $\Delta(D - \sigma) + \sigma$, where Δ is the uncorrected density of the body, D is the density of the water, and σ is the density of the air. The table below gives the correction to be applied to Δ . D is taken as .9992 (correct to 1 part in 2000 between 10° and 18° C., see p. 22) and σ as .0012 (see p. 25). — means that the correction has to be subtracted from Δ . (See Stewart and Gee, "Practical Physics," vol. i.)

Δ	Corr.	Δ	Corr.	Δ	Corr.	Δ	Corr.	Δ	Corr.	Δ	Corr.
0.5	+ .0002	4.0	— .0068	7.5	— .0138	8.4	— .0156	9.5	— .0178	18.0	— .0308
1.0	— .0008	4.5	— .0078	7.8	— .0144	8.5	— .0158	10.0	— .0188	17.0	— .0328
1.5	— .0018	5.0	— .0088	7.9	— .0146	8.6	— .0160	11.0	— .0208	18.0	— .0348
2.0	— .0028	5.5	— .0098	8.0	— .0148	8.7	— .0162	12.0	— .0228	19.0	— .0368
2.5	— .0038	6.0	— .0108	8.1	— .0150	8.8	— .0164	13.0	— .0248	20.0	— .0388
3.0	— .0048	6.5	— .0118	8.2	— .0152	8.9	— .0166	14.0	— .0268	21.0	— .0408
3.5	— .0058	7.0	— .0128	8.3	— .0154	9.0	— .0168	15.0	— .0288	22.0	— .0428

DENSITY OF DAMP AIR

The density of damp air may be derived from the expression $\sigma = \sigma_d(H - 0.378p)/H$, where σ_d is the density of dry air at a pressure H mms. (see p. 25), H is the barometric height, and p is the pressure of water-vapour in the air.

HYDROMETERS

Common: Density = degrees/1000.

Baumé: Density at 15° = $144.3/(144.3 - \text{Baumé degrees})$.

Twaddell: Density = $1 + (\text{Twaddell degrees}/200)$.

Sikes: One degree = a density interval of .002 on the average.

DENSITIES

DENSITY OF WATER

In grams per millilitre.* Pure air-free water under 1 atmos. Temps. on const.-vol. H. scale. Water has a **maximum density** at 3°·98 (Chappuis, 1897; Thiesen, Scheel and Diesselhorst; De Coppet, 1903). The temp. (t_m) of maximum density at different pressures (p), measured in atmos., is given by $t_m = 3\cdot98 - \cdot0225(p - 1)$.

The **specific volume** is the reciprocal of the density. For reciprocals, see p. 136. (See Chappuis, *Trav. et Mém. Bur. Intl.*, 13, 1907.)

For density of ice see p. 21; of steam, p. 26. [* 1 litre = 1000·027 c.cs.]

Density of water at -10° = '99815; at -5° = '99930.

Temp.	0	2	4	6	8	10	12	14	16	18
0° C.	'99987	'99997	1'00000	'99997	'99988	'99973	'99953	'99927	'99897	'99862
20	'99823	'99780	'99732	'99681	'99626	'99567	'99505	'99440	'99371	'9930
40	'9922	'9915	'9907	'9898	'9890	'9881	'9872	'9862	'9853	'9843
60	'9832	'9822	'9811	'9801	'9789	'9778	'9767	'9755	'9743	'9731
80	'9718	'9706	'9693	'9680	'9667	'9653	'9640	'9626	'9612	'9598
100	'9584	—	—	—	—	'951	—	—	—	—

Density at 150° = '917; at 200° = '863; at 250° = '79; at 300° = '70.

DENSITY OF MERCURY

In grams per c.c. Hydrogen scale of temp. For reciprocals, see p. 136. (See Chappuis, *Trav. et Mém. Bur. Intl.*, 13, 1907.)

Temp.	0	2	4	6	8	10	12	14	16	18
-20° C.	¹³ '6450	¹³ '6400	¹³ '6351	¹³ '6301	¹³ '6251	¹³ '6202	¹³ '6152	¹³ '6103	¹³ '6053	¹³ '6004
0	'5955	'5905	'5856	'5806	'5757	'5708	'5659	'5609	'5560	'5511
20	'5462	'5413	'5364	'5315	'5266	'5217	'5168	'5119	'5070	'5022
40	'4973	'4924	'4875	'4826	'4778	'4729	'4680	'4632	'4583	'4534
60	'4486	'4437	'4389	'4340	'4292	'4243	'4195	'4146	'4098	'4050
80	'4001	'3953	'3904	'3856	'3808	'3759	'3711	'3663	'3615	'3566
	0	20	40	60	80	100	120	140	160	180
100	¹³ '3518	¹³ '3504	¹³ '3487	¹³ '3469	¹³ '3452	¹³ '3435	¹³ '3418	¹³ '3401	¹³ '3384	¹³ '3367
300	¹² '881	¹² '884	¹² '887	¹² '890	—	—	—	—	—	—

DENSITY OF ETHYL ALCOHOL, C₂H₅OH. Aq

In grams per c.c. % indicates grams of C₂H₅OH in 100 grams of aqueous solution. Hydrogen scale of temp. (Calculated by E. W. Morley from Mendeleeff's Observations, *Four. Am. Chem. Soc.*, Oct. 1904.)

At 17° C.

%	0	1	2	3	4	5	6	7	8	9
0	'9988	'9969	'9951	'9933	'9916	'9899	'9884	'9869	'9854	'9840
10	'9826	'9813	'9800	'9787	'9775	'9762	'9750	'9737	'9725	'9713
20	'9700	'9687	'9674	'9661	'9647	'9633	'9619	'9604	'9589	'9573
30	'9557	'9540	'9524	'9506	'9489	'9470	'9452	'9433	'9414	'9394
40	'9375	'9354	'9334	'9313	'9292	'9271	'9250	'9228	'9207	'9185
50	'9163	'9140	'9118	'9096	'9073	'9051	'9028	'9005	'8982	'8959
60	'8936	'8913	'8890	'8867	'8843	'8820	'8797	'8773	'8749	'8726
70	'8702	'8678	'8655	'8631	'8607	'8582	'8558	'8534	'8510	'8485
80	'8461	'8436	'8411	'8386	'8361	'8336	'8310	'8285	'8259	'8232
90	'8206	'8179	'8152	'8124	'8096	'8068	'8039	'8010	'7980	'7950
100	'7919	—	—	—	—	—	—	—	—	—

For other temperatures, interpolate from the above and the following:—

At 22° C.

0%, '9978; 10%, '9813; 20%, 'c678; 30%, '9526; 40%, '9338; 50%, '9122; 60%, '8895; 70%, '8660; 80%, '8417; 90%, '8162; 100%, '7876.

DENSITY OF HYDROCHLORIC ACID, HCl . AqGrams per c.c. at 15°C . (Lunge and Marchlewski, 1891.)

Dens.	Grams HCl in		Dens. Change for $\pm 1^{\circ}$.	Dens.	Grams HCl in		Dens. Change for $\pm 1^{\circ}$.	Dens.	Grams HCl in		Dens. Change for $\pm 1^{\circ}$.
	100 gm.	1 litre of Solution.			100 gm.	1 litre of Solution.			100 gm.	1 litre of Solution.	
1'01	2'14	22	'00016	1'08	16'15	174	'00035	1'15	29'6	340	'00052
1'02	4'13	42	'00019	1'09	18'1	107	'00038	1'16	31'5	366	'00054
1'03	6'15	64	'00021	1'10	20'0	200	'00040	1'17	33'5	392	'00056
1'04	8'16	85	'00024	1'11	21'9	243	'00043	1'18	35'4	418	'00058
1'05	10'17	107	'00027	1'12	23'8	267	'00045	1'19	37'2	443	'00059
1'06	12'19	129	'00030	1'13	25'7	291	'00048	1'20	39'1	469	'00060
1'07	14'17	152	'00032	1'14	27'7	315	'00050				

DENSITY OF NITRIC ACID, HNO_3 . AqGrams per c.c. at 15°C . % $\text{N}_2\text{O}_5 = 857 \times \% \text{HNO}_3$ —by weight. (Lunge and Rey, 1891.)

Dens.	Grams HNO_3 in		Dens. Change for $\pm 1^{\circ}$.	Dens.	Grams HNO_3 in		Dens. Change for $\pm 1^{\circ}$.	Dens.	Grams HNO_3 in		Dens. Change for $\pm 1^{\circ}$.
	100 gm.	1 litre of Solution.			100 gm.	1 litre of Solution.			100 gm.	1 litre of Solution.	
1'02	3'70	38	'00022	1'22	35'3	430	'00080	1'42	69'8	991	'00137
1'04	7'26	75	'00028	1'24	38'3	475	'00086	1'44	74'7	1075	'00143
1'06	10'7	113	'00034	1'26	41'3	521	'00091	1'46	80'0	1168	'00149
1'08	13'9	151	'00040	1'28	44'4	568	'00097	1'48	86'0	1274	'00154
1'10	17'1	188	'00045	1'30	47'5	617	'00103	1'50	94'1	1411	'00160
1'12	20'2	227	'00051	1'32	50'7	669	'00109	1'504	96'0	1444	'00161
1'14	23'3	266	'00057	1'34	54'1	725	'00114	1'508	97'5	1470	'00162
1'16	26'4	306	'00062	1'36	57'6	783	'00120	1'512	98'5	1490	'00163
1'18	29'4	347	'00068	1'38	61'3	846	'00126	1'516	99'2	1504	'00164
1'20	32'4	388	'00074	1'40	65'3	914	'00132	1'520	99'7	1515	'00166

DENSITY OF SULPHURIC ACID, H_2SO_4 . AqGrams per c.c. at 15°C . % $\text{SO}_3 = 816 \times \% \text{H}_2\text{SO}_4$ —by weight. (Lunge and Isler, 1895.)

Density.	Grams H_2SO_4 in		Density.	Grams H_2SO_4 in		Density.	Grams H_2SO_4 in	
	100 gm.	1 litre of Solution.		100 gm.	1 litre of Solution.		100 gm.	1 litre of Solution.
1'02	3'03	31	1'44	54'1	779	1'822	90'4	1647
1'04	5'96	62	1'46	56'0	817	1'824	90'8	1656
1'06	8'77	93	1'48	57'8	856	1'826	91'2	1666
1'08	11'60	125	1'50	59'7	896	1'828	91'7	1676
1'10	14'35	158	1'52	61'6	936	1'830	92'1	1685
1'12	17'01	191	1'54	63'4	977	1'832	92'5	1695
1'14	19'61	223	1'56	65'1	1015	1'834	93'0	1706
1'16	22'19	257	1'58	66'7	1054	1'836	93'8	1722
1'18	24'76	292	1'60	68'5	1096	1'838	94'6	1739
1'20	27'3	328	1'62	70'3	1139	1'840	95'6	1759
1'22	29'8	364	1'64	72'0	1181			
1'24	32'3	400	1'66	73'6	1222	1'8405	95'9	1765
1'26	34'6	435	1'68	75'4	1267	1'8410	97'0	1786
1'28	36'9	472	1'70	77'2	1312	1'8415	97'7	1799
1'30	39'2	510	1'72	78'9	1357	1'8410	98'2	1808
1'32	41'5	548	1'74	80'7	1404	1'8405	98'7	1816
1'34	43'7	586	1'76	82'4	1451	1'8400	99'2	1825
1'36	45'9	624	1'78	84'5	1504	1'8395	99'4	1830
1'38	48'0	662	1'80	86'9	1564	1'8390	99'7	1834
1'40	50'1	702	1'81	88'3	1598	1'8385	99'9	1838
1'42	52'1	740	1'82	90'0	1639			

DENSITIES : ALKALIES

DENSITY OF AMMONIA, NH_3 . AqGrams per c.c. at 15°C .

Dens.	Grams NH_3 in		Dens. Change for $\pm 1^\circ$	Dens.	Grams NH_3 in		Dens. Change for $\pm 1^\circ$	Dens.	Grams NH_3 in		Dens. Change for $\pm 1^\circ$
	100 gm.	1 litre of Solution.			100 gm.	1 litre of Solution.			100 gm.	1 litre of Solution.	
'996	'91	9'1	'00019	'956	11'03	105'4	'00031	'916	23'03	210'9	'00049
'992	1'84	18'2	'00020	'952	12'17	115'9	'00033	'912	24'33	221'9	'00051
'988	2'80	27'7	'00021	'948	13'31	126'2	'00035	'908	25'65	232'9	'00053
'984	3'80	37'4	'00022	'944	14'46	136'5	'00037	'904	26'98	243'9	'00055
'980	4'80	47'0	'00023	'940	15'63	146'9	'00039	'900	28'33	255'0	'00057
'976	5'80	56'6	'00024	'936	16'82	157'9	'00041	'896	29'69	266'0	'00059
'972	6'80	66'1	'00025	'932	18'03	168'1	'00042	'892	31'05	277'0	'00060
'968	7'82	75'7	'00026	'928	19'25	178'6	'00043	'888	32'50	288'6	'00062
'964	8'84	85'2	'00027	'924	20'49	189'3	'00045	'884	34'10	301'4	'00064
'960	9'91	95'1	'00029	'920	21'75	200'1	'00047	'880	35'70	314'2	'00066

DENSITY OF SODIUM HYDROXIDE, NaOH . AqGrams per c.c. at 18°C . The percentages indicate grams of NaOH in 100 grams of solution. (Bousfield and Lowry, 1905.)

%	Density.	%	Density.	%	Density.	%	Density.	%	Density.
0	'9986	10	1'1038	20	1'2202	30	1'3290	40	1'4314
1	1'0100	11	1'1208	21	1'2312	31	1'3396	41	1'4411
2	1'0213	12	1'1319	22	1'2422	32	1'3502	42	1'4508
3	1'0324	13	1'1429	23	1'2532	33	1'3605	43	1'4604
4	1'0435	14	1'1540	24	1'2641	34	1'3708	44	1'4699
5	1'0545	15	1'1650	25	1'2751	35	1'3811	45	1'4794
6	1'0656	16	1'1761	26	1'2860	36	1'3913	46	1'4890
7	1'0766	17	1'1871	27	1'2968	37	1'4014	47	1'4985
8	1'0877	18	1'1982	28	1'3076	38	1'4115	48	1'5080
9	1'0987	19	1'2092	29	1'3184	39	1'4215	49	1'5174

DENSITY OF SODIUM CARBONATE, Na_2CO_3 . AqGrams per c.c. at 15°C . (Lunge.)

Density.	Grams Na_2CO_3 in		Density.	Grams Na_2CO_3 in		Density.	Grams Na_2CO_3 in	
	100 gm.	1 litre of Solution.		100 gm.	1 litre of Solution.		100 gm.	1 litre of Solution.
1'007	'67	6'8	1'060	5'71	60'5	1'116	10'95	122'2
1'014	1'33	13'5	1'067	6'37	68'0	1'125	11'81	132'9
1'022	2'09	21'4	1'075	7'12	76'5	1'134	12'61	143'0
1'029	2'76	28'4	1'083	7'88	85'3	1'142	13'16	150'3
1'036	3'43	35'5	1'091	8'62	94'0	1'152	14'24	164'1
1'045	4'29	44'8	1'100	9'43	103'7			
1'052	4'94	52'0	1'108	10'19	112'9			

Change of density per 1°C . (0° to 30°), 0 to 7% = '0002; 11 to 20% = '0004.DENSITY OF CALCIUM CHLORIDE, CaCl_2 . AqGrams per c.c. at $17'9^\circ \text{C}$. The percentages indicate grams of anhydrous CaCl_2 in 100 grams of solution. (Pickering, 1894.)

%	Density.	%	Density.	%	Density.	%	Density.	%	Density.
1	1'007	11	1'094	21	1'189	31	1'294	41	1'406
3	1'024	13	1'112	23	1'209	33	1'316	43	1'429
5	1'041	15	1'131	25	1'229	35	1'338		
7	1'058	17	1'150	27	1'250	37	1'361		
9	1'076	19	1'169	29	1'272	39	1'384		

DENSITIES OF SOME AQUEOUS SOLUTIONS

Grams per c.c. at 18° C. The indicated % is the number of grams of anhydrous substance in 100 grams of solution. (Kohlrausch, "Prakt. Phys.")

Substance.	5%	10%	15%	20%	25%	Substance.	5%	10%	15%	20%
NaCl .	1'034	1'071	1'109	1'148	1'190	MgSO ₄ .	1'050	1'104	1'160	1'220
NaNO ₃	1'033	1'068	1'105	1'144	1'185	BaCl ₂ .	1'044	1'093	1'147	1'204
NaA .	1'025	1'051	1'078	1'105	1'132	NH ₄ Cl.	1'014	1'029	1'043	1'057
H ₃ PO ₄ .	1'027	1'054	1'083	1'114	1'145	CuSO ₄ .	1'051	1'107	1'167	1'230
ZnSO ₄ .	1'051	1'107	1'167	1'232	1'305	KCl .	1'031	1'064	1'098	1'133
FeCl ₃ .	1'130	1'175	1'226	1'278	1'331	KNO ₃ .	1'030	1'063	1'097	1'133
SrCl ₂ .	1'044	1'093	1'146	1'202	1'256	K ₂ SO ₄ .	1'039	1'081	—	—
MgCl ₂ .	1'042	1'086	1'130	1'176	1'225	K ₂ Cr ₂ O ₇	1'035	1'072	1'109	—

Substance.	5%	10%	15%	20%	25%	30%	35%	40%	45%	50%
KBr .	1'035	1'073	1'114	1'157	1'204	1'254	1'307	1'365	1'429	—
KI .	1'036	1'076	1'120	1'168	1'218	1'273	1'332	1'397	1'468	1'545
K ₂ CO ₃ .	1'044	1'091	1'140	1'191	1'244	1'299	1'356	1'415	1'477	1'541
LiCl .	1'027	1'056	1'085	1'115	1'147	1'181	1'217	1'255	—	—
CdSO ₄ .	1'049	1'103	1'161	1'224	1'295	1'372	1'457	—	—	—
AgNO ₃ .	1'042	1'089	1'140	1'196	1'255	1'321	1'394	1'477	1'570	1'674
PbA ₂ .	1'036	1'075	1'118	1'163	1'212	1'265	1'322	1'386	—	—
Sugar* .	1'018	1'039	1'060	1'081	1'104	1'128	1'152	1'177	1'203	1'230

* 60%, 1'287; [75%, 1'380 (supersaturated)].

DENSITY OF DRY AIR AT DIFFERENT TEMPERATURES AND PRESSURES

Grams per c.c.; pressures in mm. of mercury at 0° C. lat. 45°; $g = 980.62$ cms. per sec.². These densities are calculated by the expression $\frac{0.001293}{(1 + 0.00367t)} \cdot \frac{H}{760}$, where 0.001293 is due to Leduc, 1898, and Rayleigh, 1893 (p. 26); and 0.00367 to Regnault. For density of damp air, see p. 21.

Temp. (°).	Pressure in Millimetres (H).							
	710	720	730	740	750	760	770	780
0° C.	0.001208	0.001225	0.001242	0.001259	0.001276	0.001293	0.001310	0.001327
2	0.001199	0.001216	0.001233	0.001250	0.001267	0.001284	0.001300	0.001317
4	0.001190	0.001207	0.001224	0.001241	0.001258	0.001274	0.001291	0.001308
6	0.001182	0.001199	0.001215	0.001232	0.001248	0.001265	0.001282	0.001298
8	0.001173	0.001190	0.001207	0.001223	0.001240	0.001256	0.001273	0.001289
10	0.001165	0.001182	0.001198	0.001214	0.001231	0.001247	0.001264	0.001280
12	0.001157	0.001173	0.001190	0.001206	0.001222	0.001238	0.001255	0.001271
14	0.001149	0.001165	0.001181	0.001197	0.001214	0.001230	0.001246	0.001262
16	0.001141	0.001157	0.001173	0.001189	0.001205	0.001221	0.001237	0.001253
18	0.001133	0.001149	0.001165	0.001181	0.001197	0.001213	0.001229	0.001245
20	0.001125	0.001141	0.001157	0.001173	0.001189	0.001205	0.001220	0.001236
22	0.001118	0.001133	0.001149	0.001165	0.001181	0.001196	0.001212	0.001228
24	0.001110	0.001126	0.001141	0.001157	0.001173	0.001188	0.001204	0.001220
26	0.001103	0.001118	0.001134	0.001149	0.001165	0.001180	0.001196	0.001211
28	0.001095	0.001111	0.001126	0.001142	0.001157	0.001173	0.001188	0.001203
30	0.001088	0.001103	0.001119	0.001134	0.001149	0.001165	0.001180	0.001195

GASEOUS DENSITIES

DENSITIES OF GASES

Only those gases for which accurate density determinations have been made are included in this table (see also p. 10). Other gases will be found in the table below. For density of air under different temperatures and pressures, see p. 25.

Densities are in grams per litre (1000/027 c.c.s.; see p. 10) at 0° C. under 760 mm. of mercury at 0° C. and lat. 45° ($g = 980.62$), i.e. under a pressure of 1.01323×10^6 dynes per sq. cm. (After P. A. Guye, *Chem. News*, 1908.)

Gas.	Density and Observer.	Accepted density.	Density rel. to O
Air	1.2927 L.; 1.2928 R.	Grams/litre. 1.2928	0.90469
Oxygen, O ₂	{ 1.4288 L.; 1.42905 R.; 1.42900 M.; 1.42896 Gr.; 1.4292 J.P.	1.42900	1.00000
Hydrogen, H ₂	0.08982 L.; 0.08998 R.; 0.089873 M.	0.08987	0.06289
Nitrogen, N ₂	1.2503 L.; 1.2507 R.; 1.2507 Gr.	1.2507	0.87523
Argon, A	1.7809 R.; 1.7808 Ra.	1.7809	1.2463
Nitrous oxide, N ₂ O	1.9780 L.; 1.9777 R.; 1.9774 G.P.	1.9777	1.3840
Nitric oxide, NO	1.3429 L.; 1.3402 Gr.; 1.3402 G.D.	1.3402	0.93786
Ammonia, NH ₃	0.7719 L.; 0.77085 P.D.; 0.7708 G.P.	0.7708	0.5394
Carbon monoxide, CO	1.2501 L.; 1.2504 R.	1.2504	0.87502
Carbon dioxide, CO ₂	1.9763 L.; 1.9769 R.; 1.9768 G.P.	1.9768	1.3833
Hydrochloric acid, HCl	1.6407 L.; 1.6397 Gr.; 1.6398 G.G.	1.6398	1.1475
Sulphur dioxide, SO ₂	2.9266 L.; 2.9266 J.P.; 2.9266 B.	2.9266	2.0480

B., Berthelot; G.D., Guye & Davila; G.G., Guye & Gazarian; G.P., Guye & Pintza; Gr., Gray; J.P., Jacqueroed & Pintza; L., Leduc; M., Morley; P.D., Perman & Davies; R., Rayleigh; Ra., Ramsay.

The densities below are all **experimental values**, and are relative to that of oxygen (O₂ = 16) at 0° and 760 mms. at lat. 45° (see above).

Gas.	Rel. dens.	Gas.	Rel. dens.	Gas.	Rel. dens.
Acetylene, C ₂ H ₂	13.32	Helium, He	1.98	Nitrogen oxychloride, NOCl	33.45
Arsine, AsH ₃	39.02	Hydrobromic acid, HBr	39.24	Nitrogen peroxide—(N ₂ O ₄) 26° 7 C.	38.37
Boron fluoride, BF ₃	33.48	Hydrofluoric acid, HF	10.32	" " 39° 8	35.62
Bromine, Br ₂ , 228° C.	79.99	Hydriodic acid, HI	63.36	" " 60° 2	30.12
Butane, C ₄ H ₁₀	29.10	Hydrogen selenide, H ₂ Se	40.47	" " 80° 6	26.06
Carbon oxychloride, COCl ₂	50.75	" sulphide, H ₂ S	17.22	" " 100° 1	24.33
" oxysulphide, COS	30.47	" telluride, H ₂ Te	65.00	" " 121° 5	23.46
Chlorine, Cl ₂	36.07	Krypton, Kr	41.5	" (NO ₂) 154° 0	22.88
" monoxide, Cl ₂ O	43.54	Methane, CH ₄ (1909)	8.03	" " 183° 2	22.73
" dioxide, ClO ₂	33.74	Methylamine, CH ₃ NH ₂	15.64	Phosphine, PH ₃	17.58
Cyanogen, C ₂ N ₂	26.16	Methyl chloride, CH ₃ Cl	25.06	Phosphorus chlorofluoride, PCl ₂ F ₃	78.19
Ethane, C ₂ H ₆	15.57	Methyl ether, C ₂ H ₆ O	23.41	" oxyfluoride, PO ₂ F ₃	53.29
Ethylamine, C ₂ H ₅ NH ₂	22.77	" fluoride, CH ₃ F	17.67	" pentafluoride, PF ₅	65.01
Ethyl chloride, C ₂ H ₅ Cl	32.13	Methylene fluoride, CH ₂ F ₂	26.21	" trifluoride, PF ₃	43.76
Ethyl fluoride, C ₂ H ₅ F	24.62	Neon, Ne (1910)	10.82	Propylene, C ₃ H ₆	21.69
Ethylene, C ₂ H ₄	14.27			Silicon fluoride, SiF ₄	52.13
Fluorine, F ₂	18.97			Xenon, Xe	65.35

DENSITY OF SATURATED WATER VAPOUR

Densities in grams per litre under different pressures. (Zeuner, 1890.)

Atmos.	0	0.5	1	1.5	2	2.5	3	3.5	4	4.5
0	—	0.315	0.606	0.887	1.16	1.43	1.70	1.97	2.23	2.49
5	2.75	3.01	3.26	3.52	3.77	4.02	4.27	4.52	4.77	5.02
10	5.27	5.52	5.76	6.01	6.25	6.50	6.74	6.99	7.23	—

ELASTICITIES

Young's Modulus, or Longitudinal Elasticity, E in dynes per sq. cm.

Rigidity, Torsion Modulus, or Shear Modulus, n in dynes per sq. cm.

Volume Elasticity, Cubic Elasticity, or Bulk Modulus, k in dynes per sq. cm.

Compressibility (cubic), $C = 1/k$.

Poisson's Ratio, σ = lateral contraction per unit breadth/longitudinal extension per unit length. For a homogeneous isotropic substance—

$$n = \frac{E}{2(1+\sigma)} \dots (a); \quad \sigma = \frac{E}{2n} - 1 \dots (b); \quad k = \frac{E}{3(1-2\sigma)} \dots (c)$$

For an isotropic solid Poisson's Ratio must lie between $+\frac{1}{2}$ and -1 , but for some materials it may, when deduced from E and n , exceed $+1$. (See Searle's "Elasticity.")

1 megabar = 10^6 dynes per sq. cm. = .987 atmos. = $1/1.013$ atmos. = the pressure measured by 750.15 mms. of mercury at 0° C. sea-level, and latitude $45^\circ = 749.66$ mms. at 0° in London.

The elasticities of a substance depend considerably upon its history. The extent of the agreement between the calculated and observed values of n and of σ below gives an indication of the degree of isotropy of the metals used. (Grüneisen, Reichsanstalt, *Ann. d. Phys.*, 1908.)

ELASTICITIES OF METALS

Metal at 18° C. (see also below and pp. 28, 29).	Young's Modulus, E .	Rigidity, n .		Poisson's Ratio, σ .		Vol. Elast. k .	Compressy C. per megabar (calculated).
	By static method or longl. vibns.	By oscilln. method.	Calcd. by formula (a).	Ob- served.	Calcd. by for- mula (b).	Calcd. by formula (c).	
Aluminium (W) *.	7.05×10^{11}	2.67×10^{11}	2.63×10^{11}	.339	.310	7.46×10^{11}	1.33×10^{-6}
Bismuth (C), pure.	3.19	—	1.20	.33	—	3.14	3.2
Cadmium (C), pure	4.99	—	1.92	.30	—	4.12	2.4
Copper (W), pure.	12.3	4.55	4.55	.337	.356	13.1	.74
Gold (W), pure.	8.0	2.77	2.80	.422	.495	16.6	.60
Iron (W), 1% C.	21.3	—	8.31	.280	—	16.1	.63
Steel (W), 1% C.	20.9	8.12	8.12	.287	.287	16.4	.62
Lead (C), pure.	1.62	—	.562	.446	—	5.00	2.0
Nickel (W) †.	20.2	—	7.70	.309	—	17.6	.57
Palladium (C), pure	11.3	5.11	4.04	.393	.101	17.6	.57
Platinum (C), pure	16.8	6.10	6.04	.387	.368	24.7	.41
Silver (W), pure.	7.90	2.87	2.86	.379	.369	10.9	.92
Tin (C), pure.	5.43	—	2.04	.33	—	5.29	1.9
Bronze (C) ‡.	8.08	3.43	2.97	.358	.177	9.52	1.05
Constantan (W) §.	16.3	6.11	6.11	.325	.329	15.5	.65
Manganin (W) .	12.4	4.65	4.65	.329	.329	12.1	.83

(C) means cast; (W) worked. * .5% Fe, .4% Cu. † 97% Ni, 1.4% Co, 1% Mn.

‡ 85.7% Cu, 7.2% Zn, 6.4% Sn. § 60% Cu, 40% Ni. || 84% Cu, 12% Mn, 4% Ni.

The (experimental) results below are mostly for **ordinary laboratory materials**, chiefly wires.

Substance.	Young's Modulus, E .	Rigidity, n .	Volume Elast. k .	Poisson's Ratio, σ .
Copper	$12.4-12.9 \times 10^{11}$ S.	$3.9-4 \times 10^{11}$ S.	14.3×10^{11} M.	.26 S.
Iron (wrought)	19-20	7.7-8.3	14.6	.27
" (cast)	10-13 G.	3.5-5.3	9.6	.23-.31
Steel	19.5-20.6	7.9-8.9	18.1 M.	.25-.33
Zinc (1% Pb)	8.7 §	3.8	—	.21
Brass (c. 66 Cu, 34 Zn).	9.7-10.2	c. 3.5	10.65 M.	.34-.40
German silver *	11.6	4.3-4.7	—	.37
Platinoid †	13.6	3.60	—	.37
Phosphor bronze ‡ . . .	12.0	4.36	—	.38 S.
Quartz fibre	5.18	3.0	1.4 H.	—
Indiarubber048-.052	.00016	—	.46-.49 Sc.
Jena Glasses, Crowns .	6.5-7.8	2.6-3.2	4.0-5.9	.20-.27
" " Flint	5.0-6.0	2.0-2.5	3.6-3.8	.22-.26

(G.) Grüneisen, 1907.

(H.) Horton, 1905.

(M.) Mallock, 1905.

(S.) Searle, 1900.

(Sc.) Schiller, 1906.

* 60 Cu, 15 Ni, 25 Zn.

† German silver with a little tungsten.

‡ 92.5 Cu, 7 Sn, .5 P.

§ Pure Zn, 12.5×10^{11} dynes/cm².

TENSILE STRENGTHS

ELASTICITIES (*contd.*)

Substance.	Young's Modulus, E, dynes/cm. ²	Temperature coefficient α in $\text{Elast}_t = \text{Elast}_{15}\{1 - \alpha(t - 15)\}$			Compressibility C. per megabar (i.e. 10^6 dynes/cm. ²) (Buchanan, <i>Proc. R. Soc.</i> , 1904).	
		At 15° C.	α for E.*	α for π †	7-11°C.; 200-300 megabars (see also pp. 27, 29).	
Iridium	5.2×10^{11} (G.)	Aluminium	21.3×10^{-4}	13.5×10^{-4}	Aluminium	1.7×10^{-6}
Rhodium	3.2 (G.)	Copper	3.64	4.0	Copper	.88
Tantalum	18.6 (Bo.)	Gold	4.8	3.3	Gold	.80
Invar	14.1	Iron	2.3	7.3	Lead	2.8 (A.)
90Pt, 10Ir	21.0	Steel	2.4	2.6	Magnesium	3.2
Silk fibre	.65 ‡	Platinum	.98	1.0	Platinum	.56
Spider thread.	.3 (B.) §	Silver	.75	4.5	Flint glass	3.0
Catgut	.32	Tin	—	5.9	Germ. glass tubing	2.57
Ice (-2°)	.28	Brass	.37	4.6	Steel	.51 (Br.)
Quartz (crystal)	6.8	German silver	. . .	6.5		
Marble	2.6	Phosphor-bronze	. . .	6.3		
Oak	1.3	Quartz fibre	—	-1.2		
Deal	.9					
Mahogany	.88					
Teak	1.66					

(A.) Amagat. (B.) Benton, 1907 and 1908. (Bo.) v. Bolton, 1905. (Br.) Bridgman, 1909. (G.) Grüneisen, 1907. * Wassmuth, 1906, and Schaefer, 1902. † Horton, 1904 and 1905. ‡ Diminishes rapidly with increasing load. § Shows marked elastic fatigue. || Pure.

TENSILE STRENGTHS OF MATERIALS

Tenacities or breaking stresses in dynes per sq. cm. The elastic limit is always exceeded before the breaking stress is reached. The process of drawing into wire seems to strengthen the material, and the finer the wire the greater is the breaking stress. (See Poynting and Thomson's "Properties of Matter.")

For crushing and shearing strengths, see Ewing's "Strength of Materials" or one of the Engineering "Pocket-books." For bursting strengths of tubing, see p. 39; for tensile strengths of liquids, see p. 39.

To reduce to kilogrammes per sq. mm., it is sufficient to divide by 10^8 ; to lbs. per sq. inch, divide by 7×10^4 ; to tons per sq. inch, divide by 1.5×10^8 . * Along the grain.

Substance.	Tenacity. dynes/cm. ²	Substance.	Tenacity. dynes/cm. ²
Aluminium, cast	6.9×10^9	White or yellow pine *	2.5×10^9
" rolled	9-15	Leather belt	c. 3
Copper, cast	1.2-1.9	Hemp rope	6-10
" rolled	2.0-2.5	Catgut	4.2
Iron, (a) cast	8-23	Spider thread	1.8
(b) wrought	2.9-4.5	Silk fibre	2.6
(c) steel castings	2.3-7.0	Quartz fibre	c. 10
Mild steel (2% C)	4.3-4.9	WIRES.	
High carbon) annld.	7.0-7.7	Aluminium	1.7-2.0
(for springs) / temprd.	9.3-10.8	Copper, hard drawn	4.0-4.6
Tungsten or chrome	11-12	" annealed	2.8-3.1
Ni steel, 5%; 12%	6.2; 14	Gold	2.6
Lead	c. 16	Iron (charcoal), hard drawn	5.4-6.2
Tin	16-38	" annealed	c. 4.6
Zinc, rolled	1.1-1.5	Steel; (1) ordinary; (2) tempd.	c. 11; 15.5
Brass (ordinary), (66 Cu) cast	1.5-1.9	" pianoforte	18.6-23.3
" (34 Zn) rolled	2.3-3.7	Nickel	5.3
Phosphor-bronze	2.5-2.8	Platinum	3.3
Gun-metal (90 Cu, 10 Sn)	1.9-2.6	Silver	2.9
Soft solder	c. 5	Tantalum	4.2
Glass	3-9	Brass	3.1-3.9
Ash, beech, oak, teak, mahogany*	6-1.1	Phosphor-bronze, hard drawn	6.9-10.8
Fir, pitch-pine*	4-8	German silver	4.6
Red or white deal*	3-7		

COMPRESSIBILITIES OF ELEMENTS

Coefficient of compressibility $C = \frac{1}{V} \cdot \frac{\delta V}{\delta p}$, where δV is the change in volume of a volume V under a change of pressure δp (temp. constant).

The values of C below are per megabar (*i.e.* 10^6 dynes per sq. cm.). To express as compressibility per atmosphere, increase C by $\frac{1}{80}$ of its value. Room temp. Pressure range, 100–500 megabars. Based on compressibility of mercury = $\cdot 0,371$ per megabar. The results show a periodic relation with atomic weight. See also pp. 27, 28. (Richards, *Zeit. Phys. Chem.*, 61, 1907, and *Journ. Chem. Soc.*, 1911.)

Element.	C	Element.	C	Element.	C	Element.	C
Al. . .	1.3×10^{-6}	Cl (liq.).	95×10^{-6}	Hg . .	3.71×10^{-6}	Si. . .	16×10^{-6}
Sb. . .	2.2 "	Cr. . .	7 "	Mo . .	26 "	Ag . .	84 "
As. . .	4.3 "	Cu . .	54 "	Ni . .	27 "	Na . .	15.4 "
Bi. . .	2.8 "	Au . .	47 "	Pd . .	38 "	S . .	12.5 "
Br. . .	51.8 "	I . .	13 "	P, red .	9.0 "	Tl . .	2.6 "
Cd . .	1.9 "	Fe . .	40 "	white .	20.3 "	Sn . .	1.7 "
Cs. . .	61 "	Pb . .	2.2 "	Pt . .	21 "	Zn . .	1.5 "
Ca. . .	5.5 "	Li . .	8.8 "	K . .	31.5 "		
C, diamond	5 "	Mg . .	2.7 "	Rb . .	40 "		
graphite	3 "	Mn . .	67 "	Se . .	11.8 "		

COMPRESSIBILITIES OF LIQUIDS

C = compressibility per megabar (*i.e.* 10^6 dynes per cm.²). To express as compressibility per atmosphere, increase C by $\frac{1}{80}$ of its value.

As the pressure increases C becomes less. In general a rise in temperature increases the compressibility of a liquid; but water, however, shows a minimum value of C at about 50° C. (Amagat). The compressibility of a solution diminishes as the concentration increases (see Poynting and Thomson's "Properties of Matter").

Where the limits of pressure are not given, they are—for Amagat, 8–37 atmos.; for Röntgen, 8 atmos.; for Richards, 100–200 atmos.

Liquid.	Temp.	Comp. C per megabar.	Liquid.	Temp.	Comp. C per megabar.
Water, 1–25 atmos. (A.)	15° C.	48.9×10^{-6}	Carbon tetrachloride		
900–1000 " (A.)	15	36.3 "	(Ri.)	20° C.	89.6×10^{-6}
900–1000 " (A.)	198	55.4 "	Carbon bisulphide (A.)	15.6	85.9 "
2500–3000 " (A.)	14.2	25.8 "	Ether, 1–50 atmos. (A.)	0	145.2 "
Sea-water (Gra-si, 1851)	—	43.1 "	900–1000 " (A.)	0	64.2 "
Mercury . . . (A.)	20	3.82 "	" " (A.)	198	142.2 "
" " " (Ri.)	15	3.71 "	Methyl acetate . (A.)	14.3	95.8 "
Methyl alcohol, CH ₃ OH			Ethyl acetate . . (A.)	13.3	102.7 "
(A.)	14.7	102.7 "	" bromide . (A.)	99.3	291.3 "
Ethyl alcohol—			" chloride . (A.)	15.2	151.1 "
1–500 atm. (A.)	0	76 "	Acetic acid, 1–16 atm.		
150–200 atm. (Ba.)	310	414.7 "	(C. & S.)	0	40.2 "
Propyl alcohol,			Glycerine, C ₃ H ₈ (OH) ₃		
C ₃ H ₇ OH . . (R.)	17.7	95.8 "	(Q.)	20.5	24.8 "
Propyl alcohol iso- (R.)	17.8	101.7 "	Olive oil . . . (Q.)	20.5	62.5 "
Butyl alcohol, C ₄ H ₉ OH			Paraffin oil (de Metz,		
(R.)	17.4	88.9 "	1890)	14.8	61.9 "
Butyl alcohol iso- (R.)	17.9	96.8 "	Petroleum (Martini) .	16.5	68.7 "
Amyl alcohol,			Pentane, C ₅ H ₁₂ . (G.)	20	314 "
C ₅ H ₁₁ OH . . (R.)	17.7	89.4 "	Benzene, C ₆ H ₆ . (R.)	17.9	90.8 "
Chloroform . . (Ri.)	20	9.4 "	Turpentine, C ₁₀ H ₁₆ (Q.)	19.7	78.14 "

(A.) Amagat, *Comptes Rendus*, 1884–93; (B.) Bartoli, 1896; (Ba.) Barus, 1891; (C. & S.), Colladon and Sturm, 1827; (G.) Grimaldi, 1886; (Q.) Quincke, *Wied. Ann.*, 19, 1883; (R.) Röntgen, *Wied. Ann.*, 44, 1891; (Ri.) Richards, 1907.

VISCOSITIES

VISCOSITIES OF LIQUIDS

If two parallel planes are at unit distance apart in a fluid, and one of them is moving in its own plane with unit velocity relatively to the other plane, then the tangential force exerted per unit area on each of the planes is equal to the viscosity. The dimensions of a viscosity are $ML^{-1}T^{-1}$.

For the capillary-tube method of determining viscosities, Poiseuille's formula is, Viscosity $\eta = \frac{\pi p r^4 t}{8 l V}$, where p is the pressure difference between the two ends of the tube, r the radius of the tube, l its length, V the volume of liquid delivered in a time t .

VISCOSITY OF WATER

Determined by an efflux method and corrected for kinetic energy of outflow. (Hosking, *Phil. Mag.*, 1909, 1, 502 ; 2, 260.)

Temp.	Viscosity.	Temp.	Viscosity.	Temp.	Viscosity.	Temp.	Viscosity.
0° C.	c.g.s. '01793	20° C.	'01006	50° C.	'00550	90° C.	'00316
5	'01522	25	'00893	60	'00469	100	'00284
10	'01311	30	'00800	70	'00406	124 *	'00223
15	'01142	40	'00657	80	'00356	153 *	'00181

* de Haas, 1894.

VISCOSITY OF MERCURY

(Koch, 1881.)

Temp.	-20° C.	0°	20°	50°	100°	200°	300°
Viscosity (c.g.s.)	'0186	'0169	'0156	'0141	'0122	'0101	'0093

VISCOSITIES OF VARIOUS LIQUIDS

Substance.	0° C.	10°	20°	30°	40°	50°	60°	70°
	c.g.s.							
Methyl alcohol, CH_4O	'00813	'00686	'00591	'00515	'00450	'00396	'00349	—
Ethyl " C_2H_6O	'0177	'0145	'0119	'00989	'00827	'00697	'00591	'00504
Propyl " C_3H_8O	'0388	'0292	'0225	'0178	'0140	'0113	'00919	'00757
Isopropyl " . . .	'0456	'0324	'0237	'0175	'0133	'0103	'00804	'00642
Ether $(C_2H_5)_2O$. . .	'00286	'00258	'00234	'00212	—	—	—	—
Chloroform, $CHCl_3$. . .	'00700	'00626	'00564	'00511	'00465	'00426	'00390	—
Carbon tetrachloride . . .	'0135	'0113	'00969	'00841	'00738	'00653	'00583	'00524
" bisulphide . . .	'00429	'00396	'00367	'00342	'00319	—	—	—
" dioxide (liq.) . . .	—	'00085	'00071	'00053	—	—	—	—
Benzene, C_6H_6 . . .	'00902	'00759	'00649	'00562	'00492	'00437	'00390	'00351
Aniline, $C_6H_5NH_2$. . .	—	'0655	'0440	'0319	'0241	'0189	'0156	—
Glycerine, $C_3H_5(OH)_3$ 46°	21°	8°	3°	—	—	—	—	—
Bromine . . .	'0126	'0111	'00993	'00898	'00817	'00746	—	—
Turpentine, dens. = .87	'0225	'0178	'0149	'0127	'0107	'00926	'00821	'00728
Pentane (n), C_5H_{12} . . .	'00283	'00255	'00232	'00212	—	—	—	—
Hexane (n), C_6H_{14} . . .	'00396	'00355	'00320	'00290	'00264	'00241	'00221	—
Formic acid, HCO_2H . . .	—	'0224	'0178	'0146	'0122	'0103	'0089	'0077
Acetic acid, CH_3CO_2H . . .	—	—	'0122	'0104	'0090	'0079	'0070	'0062
Propionic acid, $C_3H_7O_2$. . .	'0152	'0129	'0110	'0096	'0084	'0075	'0067	'0060
Butyric " $C_4H_9O_2$. . .	'0228	'0185	'0154	'0130	'0112	'0097	'0085	'0076
Isobutyric " " . . .	'0188	'0157	'0131	'0113	'0098	'0086	'0076	'0068
Methyl formate . . .	'00429	'00384	'00347	'00317	—	—	—	—
Ethyl " . . .	'00505	'00448	'00402	'00362	'00328	'00299	—	—
Methyl acetate . . .	'00478	'00425	'00381	'00344	'00312	'00284	—	—

Machine oil, $c. 1/19^\circ$; olive oil, $'99/15^\circ$; paraffin oil, $c. '02/19^\circ$; rape oil, $1'6/20^\circ$.

RELATIVE VISCOSITIES OF SOME AQUEOUS SOLUTIONS

Strength of solutions 1 normal. Viscosities relative to that of water at same temp. For a complete list, see Stöckl in L.B.M., and Moore, *Phys. Rev.*, 1895.

Substance.	Temp.	Relative Viscosity.	Substance.	Temp.	Relative Viscosity.
Ammonia	25° C.	1.02	Potassium chloride .	17° 6 C.	.98
Ammonium chloride	17.6	.98	Potassium iodide . .	17.6	.91
Calcium chloride	20	1.31	Sodium hydrate . . .	25	1.24
Hydrochloric acid .	25	1.07	Sulphuric acid . . .	25	1.09

VISCOSITIES OF SOLIDS

Venice turpentine * at 17° 3, 1300, c.g.s. Shoemaker's wax † at 8°, 4.7×10^6 c.g.s.
Pitch † at 0°, 51×10^{10} ; at 15°, 1.3×10^{10} . Soda glass † at 575°, 11×10^{12} .
Glacier ice, † 12×10^{13} . Golden Syrup (Lyle), 1400/12°.

* R. Ladenburg, 1906.

† Trouton and Andrews, 1904.

‡ Deeley, 1908.

VISCOSITIES OF GASES AND VAPOURS

Clerk Maxwell showed in 1860 that, on the basis of the kinetic theory, the coefficient of viscosity of a gas would be independent of the pressure, and would vary as the square root of the absolute temperature. The first relation is true except at very low pressures; the second deduction is not supported by experiment.

Of the formulæ connecting gaseous viscosity (η) and temperature (t), there are the convenient but only approximate relation of O. E. Meyer, $\eta_t = \eta_0 (1 + \alpha t)$, where α is a const.; and the less manageable but accurate formula of Sutherland (*Phil. Mag.*, 31, 1893), who, by taking account of the effects of molecular forces in bringing about collisions which otherwise would have been avoided, derived the

expression $\eta_t = \eta_0 \frac{273 + C}{\theta + C} \cdot \left(\frac{\theta}{273} \right)^{\frac{3}{2}}$, where θ is the absolute temperature, and C is

Sutherland's constant. The formula only holds for temps. above the critical, and for pressures such that Boyle's law is approximately obeyed. Sutherland's relation

is thus of the form (which lends itself to graphical treatment), $\theta = \frac{K\theta^{3/2}}{\eta} - C$, where

K is a constant. (See Fisher, *Phys. Rev.*, 1907, 1909 *et seq.*; O. E. Meyer's "Kinetic Theory of Gases." For a bibliography of gaseous viscosity, see Pedersen, *Phys. Rev.*, 25, 1907.) The values below are for dry gases.

Gas or Vapour.	Temp.	η .	Observer.	Gas or Vapour.	Temp.	η .	Observer.
Air . . .	-21° C.	$\times 10^{-6}$	Breitenbach	Nitrogen	0° C.	$\times 10^{-6}$	v. Obermayer
	0	164	" (1901)		11	166	" (1876)
	0	173	Hogg, 1905		54	171	"
	0	171	G. & G.* 1908	Helium .	0	190	"
	0	170	Fisher, 1909		0	189	Schultze, '01
	15	171	Markowski		15	197	"
	99.6	181	" (1904)	Neon . .	185	270	"
Hydrogen	302	221	Breitenbach	Argon . .	15	312	Rankine, '10
	-21	299	" (1901)	Krypton .	0	210	Schultze, '01
	0	82	"		15	221	"
	0	86	"		184	322	"
	15	89	"	Xenon . .	15	246	Rankine, '10
Oxygen .	99	106	"	Chlorine .	15	222	"
	302	139	"		0	129	Graham, '46
	0	187	v. Obermayer		20	147	"
	15	195	" (1876)	Water (vap.)	0	90	Puluj, 1878
Nitrogen .	54	216	"	15	97	K. & W. † 1875	
	-21	157	"		100	M. & S. § 1881	

* Grindley and Gibson.

† Kundt and Warburg.

§ Meyer and Schumann.

VISCOSITIES

VISCOSITIES OF GASES AND VAPOURS (*contd.*)

Gas or Vapour.	Temp.	η .	Observer.	Gas or Vapour.	Temp.	η .	Observer.
Mercury (vap.)	0° C.	$\times 10^{-6}$ 162 * 300 380	S. Koch, '83 " "	Carbon dioxide	99° C.	$\times 10^{-6}$ 186 268	Breitenbach " (1901)
Nitrous oxide	-21 0	125 135	v. Obermayer " (1876)	Methane, CH ₄	0 20	104 120	Graham, '46 "
Nitric oxide	100 0	183 165	" Graham, '46	Ethylene, C ₂ H ₄	-21 0	89 97	Breitenbach " (1901)
Sulphur dioxide	20 0	186 123	" "	Alcohol (vap.)	15 0	102 83	" Puluj, 1878
Sulphuret ¹ hydrogen	20 0	138 115	" "	Ether (vap.)	17 78	89 142	" "
Cyanogen .	0 20	130 95	" "		0 36	69 79	" "
Carbon monoxide	0 20	107 163	" v. Obermayer (1876)	Chloroform (vap.)	0 17.4	99 103	Breitenbach " (1901)
Carbon dioxide	-21 0 15	129 139 146	Breitenbach " (1901) " "	Benzene (vap.)	61 0 19 100	189 69 79 118	" Schumann " (1884) "

* Extrapolated.

TEMPERATURE COEFFICIENTS OF VISCOSITY

Based largely on W. J. Fisher's computations (ref. above).

Gas or Vapour.	Sutherland's Consts.		Meyer's Const. α	Gas or Vapour.	Sutherland's Consts.		Meyer's Const. α
	C	K			C	K	
Air	124	150×10^{-7}	.00273	Xenon	252	246×10^{-7}	—
Hydrogen	72	66	—	Water (vap.)	72	—	—
Oxygen	127	175	.00283	Carbon monoxide	102	135	.00269
Nitrogen	110	143	.00269	" dioxide	240	158	.00350
Helium	80	148	—	Nitrous oxide	313	172	.00345
Neon	56	220	—	Ethylene	226	106	.00350
Argon	170	207	—	Chloroform (vap.)	454	159	—
Krypton	188	240	—				

SIZE, VELOCITY, AND FREE PATH OF MOLECULES

ρ = density of gas in gms./c.c. at 0° C. N = number of molecules of gas per c.c. at 0° C. and 76 cms.

p = 1 atmos. = 1.0132×10^6 dynes/cm.²

θ = absolute temperature.

R = gas constant.

b = b of Van der Waals' equation (p. 34).

k = thermal conductivity of gas (p. 52).

c_v = specific heat at const. volume (p. 58).

η = viscosity of gas (p. 31).

σ = molecular diameter in cms.

m = mass of a single molecule (in grams).

G = square root of mean square molecular vel. (cm./sec. at 0° C.).

Ω = mean molecular velocity (cm./sec.).

L = length of mean free path in cms.

Assuming a Maxwell-Boltzmann distribution of velocities—

$$G = \sqrt{3p/(Nm)} = \sqrt{3p/\rho} = \sqrt{3R\theta}$$

$$\Omega = 4G/\sqrt{6\pi} = .921G$$

$$L = \eta/(\cdot 31\rho\Omega) = 2.02\eta/\sqrt{p\rho}$$

$$\text{Collision frequency} = \Omega/L = 5 \times 10^6 \text{ per sec. for } O_2$$

SIZE, VELOCITY, AND FREE PATH OF MOLECULES (*contd.*)

MOLECULAR SIZE

The molecular diameter σ has been calculated by the following formulæ:—

1. The **viscosity** η of a gas is a function of the size of its molecules.

$$\eta = .44\rho\Omega/(\sqrt{2}N\pi\sigma^2) \quad \therefore \sigma = \{.0912\rho G/(\eta N)\}^{\frac{1}{2}}$$

2. The **thermal conductivity**, $k = 1.6\eta c_v = .158\rho\Omega c_v/N\sigma^2$

$$\therefore \sigma = \{.146\rho G c_v/(\eta k)\}^{\frac{1}{2}}$$

3. **Van der Waals'**, $b = 2\pi N\sigma^3/3 \quad \therefore \sigma = \{3b/(\pi N)\}^{\frac{1}{3}}$

4. **Limiting density**, *i.e.* density D of densest known form. $\sigma = \{6\rho/(\pi DN)\}^{\frac{1}{3}}$

The values of ρ and η used in calculating G and L below are given on pp. 26, 31. The values of σ tabulated are mostly taken from Jeans' "Dynamical Theory of Gases," or Rudolf (*Phil. Mag.*, 1909, p. 795). Jeans takes $N = 4 \times 10^{19}$, while in the table following, the more recent value 2.75×10^{19} has been used.

Gas.	G at 0° C.	Mean free path, L.	Molecular diameter σ deduced from			
			η	k	b	Lt. $\rho [= D]$
	cm./sec.	cm.	cm.	cm.	cm.	cm.
Hydrogen, H ₂	18.39×10^4	18.3×10^{-6}	2.47×10^{-8}	2.40×10^{-8}	2.32×10^{-8}	2.92×10^{-8}
Helium, He	13.11 "	28.5 "	2.18 "	—	2.30 "	4.31 "
Nitrogen, N ₂	4.93 "	9.44 "	3.50 "	3.31 "	3.53 "	2.97 "
Oxygen, O ₂	4.61 "	9.95 "	3.39 "	3.11 "	—	2.79 "
Neon, Ne	5.61 "	19.3 "	—	—	—	—
Argon, A	4.13 "	10.0 "	3.36 "	—	2.86 "	4.43 "
Krypton, Kr	2.86 "	9.49 "	—	—	3.14 "	4.93 "
Xenon, Xe	2.28 "	5.61 "	—	—	3.42 "	4.88 "
Chlorine, Cl	3.07 "	4.57 "	4.96 "	—	—	—
Methane, CH ₄	6.48 "	7.79 "	—	—	—	—
Ethylene, C ₂ H ₄	4.88 "	5.47 "	4.55 "	4.68 "	—	5.26 "
Carbon monoxide, CO	4.93 "	9.27 "	3.50 "	3.31 "	—	—
Carbon dioxide, CO ₂	3.92 "	6.29 "	4.18 "	4.32 "	3.40 "	4.42 "
Ammonia, NH ₃	6.28 "	6.95 "	—	—	—	—
Nitrous oxide, N ₂ O	3.92 "	6.10 "	4.27 "	4.20 "	—	4.58 "
Nitric oxide, NO	4.76 "	9.06 "	3.40 "	3.40 "	—	—
Sulph. hydrogen, H ₂ S	4.44 "	5.90 "	—	—	—	—
Sulph. dioxide, SO ₂	3.22 "	4.57 "	—	—	—	—
Hydrochloric acid, HCl	4.30 "	6.86 "	—	—	—	—
Water, H ₂ O	7.08 "	7.22 "	4.09 "	—	—	3.45 "

The formulæ above assume the molecules to be spherical. Sutherland (*Phil. Mag.*, 1910), adopting his formula (see p. 31) for the variation of η with temp., obtains the following values of σ . Unit, 10^{-8} cm.

H	He	A	O ₂	N ₂	N ₂ O	NO	CO	CO ₂	C ₂ H ₄	Cl ₂
2.17	1.92	2.66	2.71	2.95	3.33	2.59	2.74	2.90	3.31	3.76

CRITICAL DATA

CRITICAL DATA AND VAN DER WAALS' CONSTANTS

Critical temperature, θ_c , is the highest temperature at which a gas can be liquefied by subjecting it to pressure.

Critical pressure, p_c , is the pressure (of gas and liquid) at the critical temperature.

Critical volume, v_c , is here defined as the ratio of the volume that a gas has at the critical temp. and press. to that which it would have at 0° C. and 760 mms., i.e. it is the volume of gas at θ_c and p_c which at N.T.P. would have unit volume. Some writers take the critical volume to be the specific volume (c.c.s. per gram) at θ_c and p_c .

Most of the characteristic equations of state which have been proposed for gases take the form $(p + a/v^2)(v - b) = R\theta$, where p is the pressure, v the volume, θ the absolute temperature of the gas, and R is the "gas constant." a expresses the mutual attraction of the molecules. The "covolume" b is proportional to the space occupied by the molecules: O. E. Meyer takes $b = 4\sqrt{2}$ (volume of molecule). Van der Waals assumes a is constant: if this were true the constant volume and thermodynamic scales of temperatures would agree—they do not, however (see p. 44). Joule and Thomson, Clausius, Amagat, and Berthelot, among others, regard a as a function of θ (e.g. $a \propto 1/\theta$), and b as constant.

Assuming with Van der Waals that a and b are constants, the equation can be regarded as a cubic in v , which has its three roots equal at the critical point, whence $a = 27R^2\theta_c^2/(64p_c)$, and $b = R\theta_c/(8p_c)$.

Taking pressures in atmos., and the volume of the gas at 0° C. and 1 atmos. as 1, $R = pv/\theta = 1/273$. In these units, b is in terms of the volume of the gas at 0° C. and 1 atmos.

Example.—For CO_2 $p_c = 73$ atmos. and $\theta_c = 273 + 31.1 = 304.1$, whence $b = 304.1/(8 \times 273 \times 73) = .00191$ of the volume of the gas at 0° C. and 1 atmos.

See Preston's "Heat," Nernst's "Theoretical Chemistry," Young's "Stoichiometry," Berthelot (*Trav. et Mém. Bur. Intl.*, 1907). * Indicates calculated values.

Substance.	Critical			Van der Waals'		Observer.
	Temp. θ_c	Press. p	Vol. v_c	a.	b.	
		atmos.				
Hydrogen	-234.5° C.	20	.00264*	.00042	.00088	Olszewski, '95
Oxygen	-118	50	.00426*	.00273	.00142	v. Wroblewski, '85
Nitrogen	-146	33	.00517*	.00259	.00165	"
Air	-140	39	.00468*	.00257	.00156	Olszewski, '84
Helium	-268	2.3	.00299*	.0000615	.000995	Onnes, 1908
Neon	< -210					
Argon	-117.4	52.9	.00404*	.00259	.00135	Ramsay and
Krypton	-62.5	54.3	.00532*	.00462	.00178	Travers, 1900
Xenon	14.7	57.2	.0069*	.00818	.00230	
Chlorine	146	93.5	.00615*	.01063	.00205	Knietch, '90
Bromine	302	131*	.00605	.01434	.00202	Nadejdine, '85
Water	365	194.6	.00386*	.0118	.00150	Battelli, '90
Hydrochloric acid	52.3	86	.0052*	.00697	.00173	Dewar, 1884
Carbon monoxide	-141.1	35.9	.00505*	.00275	.00168	v. Wroblewski, '83
Carbon dioxide	31.1	73	.0066	.00717	.00191	Andrews, 1869
Carbon bisulphide	273	72.9	.0090	.02316	.00343	Battelli, 1890
Ammonia, NH_3	130	115.0	.00481*	.00798	.00161	Dewar, 1884
Nitrous oxide, N_2O	38.8	77.5	.00436	.00710	.00184	Villard, 1894
Nitric oxide, NO	-93.5	71.2	.00347*	.00257	.00116	Olszewski, '85
Nitrogen tetroxide, NO_2	171.2	147*	.00413	.00756	.00138	Nadejdine, '85
Sulphuretted hydrogen	100	88.7	.00578*	.00888	.00193	Olszewski, '90
Sulphur dioxide	155.4	78.9	.00745*	.01316	.00249	Sajotschewsky, '78
Methane, CH_4	-95.5	50	.00488*	.00357	.00162	Dewar, 1884
Acetylene, C_2H_2	36.5	61.6	.0069*	.00882	.00230	Mackintosh, '07
Ethylene, C_2H_4	10	51.7	.00752*	.00877	.00251	Olszewski, '95
Ethane, C_2H_6	34	50.2	.00839*	.01060	.0028	" [86]
Ethylalcohol, $\text{C}_2\text{H}_5\text{OH}$	243	62.7	.0071	.02407	.00377	Ramsay & Young,
Ether (C_2H_5) ₂ O	197	35.8	.0158	.03496	.00602	Battelli, '92
Chloroform, CHCl_3	260	54.9	.0133	.0293	.00445	Sajotschewsky, '78
Aniline, $\text{C}_6\text{H}_5\text{NH}_2$	425.6	52.3	.0183*	.05282	.00611	Guye & Mallet, '02
Benzene, C_6H_6	288.5	47.9	.0161*	.03726	.00537	Young, 1900

DIFFUSION OF GASES

The Coefficient of diffusion, D, is the mass of the "diffusing" gas which crosses unit area in unit time under unit concentration gradient: the dimensions of the coefficient are $\text{cm}^2 \text{sec}^{-1}$. D is inversely proportional to the total pressure of the two gases, and roughly proportional to the square of their absolute temperature. Total pressure 1 atmosphere. $\text{H}_2\text{--O}_2$ implies that H_2 is diffusing into O_2 .

(See Meyer's "Kinetic Theory of Gases.")

Gases.	$t^\circ \text{C.}$	D	Gases.	$t^\circ \text{C.}$	D	Gas (Winkelmann).	$t^\circ \text{C.}$	D into		
								Air.	CO_2	H_2
$\text{H}_2\text{--O}_2$	0°	·677, O.	CO--H_2	0°	·642, L.	Formic acid	0°	·131	·088	·513
$\text{H}_2\text{--O}_2$	0	·681, O.	$\text{CO--C}_2\text{H}_4$	0	·101, O.	Acetic	0	·106	·071	·404
$\text{H}_2\text{--ClH}_4$	0	·625, O.				Propionic acid	0	·082	·058	·326
$\text{H}_2\text{--CO}$	0	·649, O.	$\text{CO}_2\text{--CO}$	0	·131, O.	Butyric acid	0	·053	·037	·201
$\text{H}_2\text{--CO}_2$	0	·538, O.	$\text{CO}_2\text{--CO}$	0	·141, L.	Isobutyric acid	0	·07	·047	·271
$\text{H}_2\text{--C}_2\text{H}_4$	0	·483, O.	$\text{CO}_2\text{--Air}$	0	·142, L.	Me. alcohol	0	·132	·088	·500
$\text{H}_2\text{--N}_2\text{O}$	0	·535, O.	$\text{CO}_2\text{--CH}_4$	0	·146, O.; ·16, L.	Et.	0	·102	·068	·378
			$\text{CO}_2\text{--O}_2$	0	·18, L.	Propyl alcohol	0	·080	·058	·315
$\text{O}_2\text{--N}_2$	0	·171, O.	$\text{CO}_2\text{--N}_2\text{O}$	0	·1, L.; ·15, O.	Butyl	0	·068	·048	·272
$\text{O}_2\text{--H}_2$	0	·722, L.	$\text{CO}_2\text{--H}_2$	0	·55, L.	"	99	·126	·088	·504
$\text{H}_2\text{O--CO}_2$	18	·155, G.	Air--O_2	0	·178, O.	Benzene	0	·075	·053	·294
$\text{H}_2\text{O--Air}$	8	·239, G.	Air--H_2	17	·66, Sc.	Me. acetate	0	·084	·056	·328
$\text{H}_2\text{O--Air}$	15	·246, G.				Et. formate	0	·085	·057	·336
$\text{H}_2\text{O--Air}$	18	·248, G.	$\text{CS}_2\text{--Air}$	0	·1, S.	Et. acetate	0	·071	·049	·273
$\text{H}_2\text{O--Air}$	0	·203, H.				Et. butyrate	0	·057	·041	·224
						Et. iso-butyrate	0	·055	·040	·224

G., Guglielmo, 1884; H., Houdaille, 1896; L., Loeschmidt, 1870; O., v. Obermayer, 1887; S., Stefan, 1879; Sc., Schulze, 1897.

DETERMINATION OF ALTITUDES BY THE BAROMETER

Babinet's formula (*Compt. Rend.*, 1850) is, $\text{Altitude} = \frac{C(\text{H}_1 - \text{H}_2)}{\text{H}_1 + \text{H}_2}$, where H_1 = barometer reading at lower station, H_2 at upper station. If altitudes are in metres, and barometric heights in mms.,

$$C = 32(500 + t_1 + t_2)$$

where t_1 and t_2 are the corresponding station temperatures ($^\circ \text{C.}$).

In the table below the mean temperature, $(t_1 + t_2)/2$, is taken as 10°C. , and the barometric height at sea-level as 760 mm., so that altitudes are in metres above sea-level. The values are of course only approximate. Babinet's formula is not applicable to very great altitudes.

Altitude	0	100	200	300	400	500	600	700	800	900
metres.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.
0	760	751	742	733	724	716	707	699	690	682
1000	674	666	658	650	642	635	627	620	612	605

THICKNESS OF THIN METAL FOIL

Approximate thickness of the thinnest beaten metal leaf at present commercially obtainable. Unit 10^{-6} cm.

Metal	Al	Cu	Au	Pt	Ag	Dutch metal.	(Cigarette paper.)
Thickness	20	34	8	25	21	70	2500

SURFACE TENSIONS

SURFACE TENSIONS

In dynes per cm. (A) indicates liquid in contact with air, (V) indicates liquid in contact with its vapour. The surface tension of a liquid varies somewhat with the age (and contamination) of the surface.

Temperature variation. It follows from Eötvös' rule, that the surface tension T at temp. t is approximately proportional to $(t_c - t)$, where t_c is the critical temp., the constant of proportionality being much the same for chemically similar substances. The surface tension at t_c is zero. (For critical temps. see p. 34.)

See Poynting and Thomson's "Properties of Matter."

WATER ($t_c = 365^\circ \text{C.}$)

Surf. Tens. T at 15°C.	Method.	Observer.	Temp. (t).	T_t/T_{15}	Temp. (t).	T_t/T_{15}
dynes per cm.						
72.8 (A)	Vibrating jet	Bohr., <i>Phil. Trans.</i> , '09	0°C.	1.030	60°C.	.901
74.3 (A)	Vibrating jet	Pedersen, <i>P. Trans.</i> , '07	10	1.010	70	.876
74.2 (A)	Capillary waves	Kalähne, <i>Ann. d. Phys.</i> , '02	15	1.000	80	.851
73.8 (A)	Hanging drop	Sentis, 1897	20	.990	90	.827
73.3 (A)	Tension of film	Hall, 1893	30	.970	100	.80
74.3 (A)	Capillary waves	Rayleigh, <i>Phil. Mag.</i> , '93	40	.947	120	.75
73.3 (A)	Capillary tube	Volkman, 1895	50	.925	140	.70
71.4 (V)	Capillary tube	Ramsay & Shields, '93	Ramsay & Shields, '93; Volkman & Brunner			
77.6 (A)	Pull on ring	Weinberg, 1892				

Substance.		Temp. (t).	Surf. Tens.	Method.	Observer.
INORGANIC.					
			dynes cm.		
Cadmium	CO ₂	Molten	693	Weight of drop	Quincke
Gold	A	1070°C.	612	Curvature of drop	Heydweiller, '98
Lead	CO ₂	335	473	Capillary waves	Grunmach
Mercury ($T_t = T_0 - 379t$)	A	17.5	547	Capillary tube	Quincke
Potassium	CO ₂	58	364	Weight of drop	"
Sodium	CO ₂	90	520	"	"
Sulphur (M.P. 115°)	A	160	59	Press. reqd. to bubble air from cap. tube thro' liquid	Zickendraht, '06; and Quincke, '08
" (B.P.)	A	250	118		
"	A	445	44	"	"
Liquid oxygen	A	-183	13.1	Capillary waves	Grunmach, 1906
" nitrogen	A	-196	8.5	" "	" 1906
" nitrous oxide	A	-89.4	26.3	" "	" 1904
Nickelcarbonyl, Ni(CO) ₄	V	19.8	14.2	Capillary tube	Ramsay and Shields, 1893
Ammonia soln. ($d = .96$)	A	15	64.7	Vibrating jet	Pedersen, 1907
Sulph' acid sol. ($d = 1.14$)	A	15	74.4	" "	" 1907
Other solns. (see below)					
CARBON COMPOUNDS.					
Acetone, (CH ₃) ₂ CO	V	16.8	23.3	Capillary tube	{ Ramsay and Shields, 1893
	V	78.3	15.9	" "	
Acetic acid, CH ₃ CO ₂ H	V	20	23.5	" "	" "
	V	300	1.16	" "	" "
Alcohol—methyl, CH ₄ O	V	20	23	" "	" "
	V	200	5.2	" "	" "
—ethyl, C ₂ H ₅ OH	V	20	22.0	" "	" "
($T_t = T_0 - .092t$)	V	150	9.5	" "	" "
—propyl (n), C ₃ H ₇ OH	V	16.4	23.8	" "	" "
	V	78.3	18.7	" "	" "
Aniline, C ₆ H ₅ .NH ₂	A	15	43.0	Vibrating jet	Pedersen, 1907
Benzene, C ₆ H ₆	A	17.5	29.2	Capillary tube	Volkman
($T_t = T_0 - .146t$)					

SURFACE TENSIONS

Substance.		Temp. (t).	Surf. Tens.	Method.	Observer.
CARBON COMPOUNDS.— (contd.)			dynes cm.		
Butyric acid, $C_3H_7CO_2H$	V	15° C.	26.7	Capillary tube	{ Ramsay and Shields, 1893
	V	132	16.4	" "	
Carbon bisulphide . .	V	19.4	33.6	" "	" "
	V	46.1	29.4	" "	" "
Carbon tetrachloride. .	V	20	25.7	" "	" "
	V	250	1.93	" "	" "
Chloroform, $CHCl_3$. .	A	15	27.2	" "	Kaye, 1905
Ether (ethyl), $(C_2H_5)_2O$.	V	20	16.5	" "	Jaeger, 1892
($T_t = T_0 - .115t$) . .	V	150	2.9	" "	" "
Ethyl acetate, $CH_3CO_2C_2H_5$	V	20	23.6	" "	" "
	V	100	14	" "	" "
Formic acid, $HCOOH$.	V	17	37.5	" "	{ Ramsay and Shields, 1893
	V	80	30.8	" "	
Olive oil ($d/20^\circ = .91$) .	A	20	32	Curvature of drop	Magie, 1888
Paraffin oil ($d = .847$) .	A	25	26.4	Capillary tube	Frankenheim, '47
Propionic acid, $C_3H_6O_2$	V	16.6	26.6	" "	{ Ramsay and Shields, 1893
	V	132	15.5	" "	
Pyridine, C_5H_5N . . .	V	17.5	36.7	" "	{ Dutoit and Fri- derich, 1900
	V	91	26.5	" "	
Toluene, $C_6H_5 \cdot CH_3$.	A	15	28.8	Vibrating jet	Pedersen, 1907
Turpentine, $C_{10}H_{16}$. .	A	15	27.3	Capillary tube	Kaye, 1905

SURF. TENSIONS OF SOLUTIONS

The surface tension of aqueous salt solutions is generally greater than that of pure water. Dorsey (*Phil. Mag.*, 1897) has shown

$$T_n = T + A \cdot n$$

T_n is the surf. tens. of a sol. of n gram — equivalents per litre, T that of water at same temp.

Salt.	A.
NaCl	1.53
KCl	1.71
$\frac{1}{2}(Na_2CO_3)$	2.00
$\frac{1}{2}(K_2CO_3)$	1.77
$\frac{1}{2}(ZnSO_4)$	1.86

SURFACE TENSIONS AT INTER-LIQUID BOUNDARIES

Liquids at 20° C.	Surface Tension T.	Observer.
	dynes/cm.	
Water-benzene	33.6	Pockels, 1899
" chloroform † . . .	29.5	Quincke
" ether	12.2	"
" olive oil ‡	20.6	"
" paraffin oil	48.3	Pockels, 1899
Mercury-water	427 *	Gouy, 1908
" alcohol §	399	Quincke
" chloroform † . . .	399	"

* Diminishes with time.

† Density = .91.

‡ Density = 1.49.

§ Density = .79.

ANGLES OF CONTACT BETWEEN GLASS AND LIQUIDS

Angles of contact vary largely with the freshness of the surfaces in contact.

Liquid.	Angle.	Observer.	Liquid.	Angle.	Observer.
Mercury	52° 40' *	Quincke	Acetic acid	20°	Magie, '88
Water	8°-9°	"	Benzene	0°	"
Water	0° †	Wilberforce	Paraffin oil	26°	"
Methyl alcohol . . .	0°	Magie, '88	Turpentine	17°	"
Ethyl alcohol	0°	"	* For freshly formed drop, 41° 5'. † Glass quite clean.		
Ether	16°	"			
Chloroform	0°	"			

The angle of contact of water against different metals varies between 3° and 11°.

SIZE OF DROPS AND THICKNESS OF LIQUID FILMS

Reference may be made to the writings of J. J. Thomson ("Conduction of Electricity through Gases"), C. T. R. Wilson, Laby (*Phil. Trans. A*, 1908), Reinold & Rücker (*Phil. Trans.*, 1886), Lord Rayleigh, and Jonhnot (*Phil. Mag.*, 1906).

HYGROMETRY

RELATIVE HUMIDITY AND DEW-POINT

Relative humidity = $\frac{[p]_t}{[p]_t^s} \cdot 100$, where $[p]_t$ is the actual pressure of water-vapour at temperature t° , and is equal to $[p]_{t_w}^s$, the saturated vapour pressure at the dew-point ($d p$); $[p]_t^s$ is the pressure of saturated vapour at t° . For a table of saturated water-vapour pressures, see p. 40. (See "Smithsonian Meteorological Tables.")

Percentage relative humidities for different dew-points and dew-point depressions are tabulated below.

Dew-point ($d p$).	Depression of dew-point = $t^\circ - (d p)^\circ$.														
	0°C.	1°	2°	3°	4°	5°	6°	7°	8°	9°	10°	12°	14°	16°	18°
- 15° C.	100	92	85	79	73	67	62	58	53	49	46	39	34	29	26
0	100	93	87	81	75	70	65	61	57	53	50	44	38	34	30
+ 10	100	94	88	82	77	72	68	64	60	56	53	47	41	37	33
20	100	94	89	83	78	74	70	66	62	58	55	49	44	39	35
30	100	94	89	84	80	75	71	68	64	61	57	52	46	42	38

WET AND DRY BULB HYGROMETER

Apjohn (1835), August (1825), and others, by making various assumptions (some of doubtful legitimacy), have derived formulæ of the type—

$$[p]_w^s - [p]_t = AH(t - t_w)[1 + B(t - t_w)]$$

where t is the temperature of the dry bulb, t_w that of the wet, $[p]_t$ is the actual pressure of water-vapour in the air (at temperature t), $[p]_w^s$ is the saturated vapour pressure of water at the temperature (t_w) of the wet bulb, H is the barometric height, and A and B are constants. (See Love & Smeal, 1911.)

The indications of this hygrometer are so dependent on its environment that for most purposes B may be taken as zero, and H as constant, say 760 mms.

If H is measured in millimetres, and temperatures in Centigrade degrees, the following values of A are suitable for the conditions mentioned:—

$A = \cdot 00068$ for moving air, as in a ventilated hygrometer.

$A = \cdot 00075$ in a Stevenson screen as used by Meteorological Office.

$A = \cdot 0008$ in open air with slight wind.

$A = \cdot 00084$ in open air with no wind.

$A = \cdot 001$ in a small closed room.

Rizzo (1897) takes $A = \cdot 00075$ and $B = -\cdot 003$, and the table below is derived by employing these values. $[p]_w^s$ can be got from the table of saturated vapour pressures on p. 40, and thus the desired vapour pressure $[p]_t$ can be determined.

VALUES OF $[p]_w^s - [p]_t$ (Rizzo)

Barom. Press. H.	Difference of temperature of dry and wet bulb thermometers ($t - t_w$).									
	1° C.	2°	3°	4°	5°	6°	7°	8°	9°	10°
mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.	mm.
770	·57	1·13	1·69	2·23	2·78	3·30	3·81	4·32	4·87	5·31
760	·56	1·12	1·67	2·20	2·74	3·25	3·76	4·27	4·75	5·24
750	·55	1·11	1·65	2·17	2·71	3·21	3·71	4·21	4·69	5·17
730	·54	1·08	1·60	2·12	2·63	3·12	3·61	4·10	4·56	5·03
700	·52	1·03	1·51	2·03	2·52	3·00	3·46	3·93	4·37	4·82
670	·50	·99	1·47	1·94	2·42	2·87	3·32	3·76	4·19	4·62
	11° C.	12°	13°	14°	15°	16°	17°	18°	19°	20°
770	5·78	6·26	6·72	7·17	7·62	8·06	8·47	8·89	9·30	9·69
760	5·71	6·18	6·63	7·08	7·52	7·95	8·36	8·77	9·18	9·56
750	5·63	6·09	6·54	6·98	7·42	7·84	8·25	8·66	9·06	9·44
730	5·48	5·93	6·37	6·79	7·22	7·63	8·03	8·43	8·82	9·18
700	5·26	5·69	6·11	6·52	6·93	7·32	7·70	8·08	8·46	8·82
670	5·03	5·44	5·84	6·24	6·63	7·01	7·37	7·73	8·08	8·43

WET AND DRY BULB HYGROMETER (*contd.*)

GLAISHER'S FACTORS

Mr. Glaisher, in 1841-5, took many thousands of observations with the wet and dry bulb hygrometer in Greenwich, India, and Toronto, and from simultaneous readings of a Daniell's hygrometer (now recognized as being an untrustworthy instrument) drew up a table of "factors."

The factor (f) at any dry-bulb reading is defined by

$$\text{depression of dew-point} = t - t_{dp} = f(t - t_w)$$

the notation being as above. Glaisher's factors are employed by the Meteorological Office and the Meteorological stations in this country. The hygrometer readings are taken in a Stevenson screen, which is essentially a box with double louvred sides.

The factors for a range of dry-bulb temperatures are tabulated below. The formula above yields the dew-point; and the saturated vapour pressure at the dew-point gives the actual vapour pressure at t° . For a table of saturated vapour pressures, see p. 40. (See "The Observers' Handbook," Meteorological Office.)

Dry Bulb Temp. (t).	0	1	2	3	4	5	6	7	8	9
- 10° C.	8.76	8.73	8.55	8.26	7.82	7.28	6.62	5.77	4.92	4.04
0	3.32	2.81	2.54	2.39	2.31	2.26	2.21	2.17	2.13	2.10
+ 10	2.06	2.02	1.99	1.95	1.92	1.89	1.87	1.85	1.83	1.81
20	1.79	1.77	1.75	1.74	1.72	1.70	1.69	1.68	1.67	1.66
30	1.65	1.64	1.63	1.62	1.61	1.60	1.59	1.58	1.57	1.56

CHEMICAL HYGROMETER

The values below are grams of water vapour contained in a cubic metre (10^6 c.c.s.) of saturated air at 760 mms. total pressure. Calculated from Regnault's observations.

Temp.	0	1	2	3	4	5	6	7	8	9
0° C.	4.84	5.18	5.54	5.92	6.33	6.76	7.22	7.70	8.21	8.76
10	9.33	9.93	10.57	11.25	11.96	12.71	13.50	14.34	15.22	16.14
20	17.12	18.14	19.22	20.35	21.54	22.80	24.11	25.49	26.93	28.45
30	30.04	31.70	33.45	35.27	37.18	39.18	41.3	43.5	45.8	48.2

TENSILE STRENGTHS OF LIQUIDS

Liquids perfectly free from air can sustain considerable tension without rupture, e.g. water can withstand a tension of 5 atmospheres, alcohol 12, and strong sulphuric acid 12 atmospheres. Extensions of volume of 0.8% for water, 1.1% for alcohol, and 1.7% for ether have been obtained. The volume elasticity (p. 29) of alcohol is the same for extension as for compression. (See Worthington, *Phil. Trans. A.*, 1892; Dixon, *Proc. Roy. Dub. Soc.*, 1909; Berthelot, *Ann. Chim. Phys.*, **30**, 1850; Poynting and Thomson's "Properties of Matter.")

BURSTING STRENGTHS OF GLASS TUBING

Bursting pressures in atmospheres for German soda glass tubing. Most glass-tubing is in a state of considerable strain, and a factor of safety of not less than two should usually be employed. (Roebuck, *Phys. Rev.*, 1909; and Onnes and Braak, *Kon. Ak. Wet.*, Amsterdam, 1908.) Ordinary boiler water-gauge glasses stand between 12 and 24 atmospheres.

Thickness of Wall.	Bore.						
	1 mm.	2	3	4	5	6	7
1 mm.	atmos.	310	280	230	220	150	190
2	570	—	340	—	330	240	220
3	560	420	460	400	—	—	230
4	—	450	—	400	310	320	280

VAPOUR PRESSURES

VAPOUR PRESSURES

Inter- and Extrapolation of Vapour Pressures.—The Kirchhoff-Rankine-Dupré formula, $\log p = A + B/\theta + C \log \theta$, where p is the vapour pressure, θ the absolute temperature, and A, B, C are constants, is accurate and convenient (e.g. see p. 41). For values of A, B, C , see Juliusburger, *Ann. d. Phys.*, p. 618, 1900.

Ramsay and Young's Method.—If two liquids, one at absolute temperature θ and the other at θ' , have the same vapour pressure, the ratio θ/θ' , when plotted against θ , gives a straight line. This method may be used to find roughly the vap. press. of a substance at any temperature when only its boiling-point is known.

Interpolation by Logarithms.—The curve of vapour pressure (p) against temp. (t) is approximately hyperbolic, and thus $\log p$ plotted against t gives a graph of slight curvature, which over 10° intervals of t may, for approximate work, be regarded as a straight line: thus the following method of interpolation:—

Example.—Required vap. press. of water at 15° , given

t	p	$\log p$	
10°	9.2	$.964$	$\frac{.964 + 1.243}{2} = 1.104 = \log 12.7$; i.e. p at $15^\circ = 12.7$, actually it is 12.8 .
20°	17.5	1.243	

VAPOUR PRESSURE OF ICE

In mms. of mercury at 0°C. ; $g = 980.62 \text{ cms. per sec.}^2$; hydrogen (const. vol.) scale of temps. (Scheel, and Heuse, Reichsanstalt *Ann. d. Phys.*, 1909.)

Temp. . .	-50°C.	-40°	-30°	-20°	-10°	-5°	-2°	0°
Vap. press.	.030 mm.	.096	.288	.784	1.963	3.022	3.885	4.579

(SATURATED) VAPOUR PRESSURE OF WATER

In mms. of mercury at 0°C. ; $g = 980.67 \text{ cms. per sec.}^2$ Thermodynamic scale of temp. (see p. 44). From -20° to 0° the observations are due to Scheel and Heuse (v. ice); from 0° to 50° , to Thiesen and Scheel; from 50° to 200° , to Holborn and Henning, Reichsanstalt (*Ann. d. Phys.*, 26, 833, 1908). For vapour pressures at temps. near 100° see also the table of boiling-points on next page.

Vap. press. at -20°C. , .960 mm.; -10° , 2.160; -5° , 3.171; -2° , 3.958; -1° , 4.258.

Temp.	0	1	2	3	4	5	6	7	8	9
0°C.	4.579	4.924	5.290	5.681	6.097	6.541	7.011	7.511	8.042	8.606
10	9.205	9.840	10.513	11.226	11.980	12.779	13.624	14.517	15.460	16.456
20	17.51	18.62	19.79	21.02	22.32	23.69	25.13	26.65	28.25	29.94
30	31.71	33.57	35.53	37.59	39.75	42.02	44.40	46.90	49.51	52.26
	0	2	4	6	8	10	12	14	16	18
40	55.13	61.30	68.05	75.43	83.50	92.30	101.9	112.3	123.6	135.9
60	149.2	163.6	179.1	195.9	214.0	233.5	254.5	277.1	301.3	327.2
80	355.1	384.9	416.7	450.8	487.1	525.8	567.1	611.0	657.7	707.3
100	760.0	815.9	875.1	937.9	1004	1074.5	1149	1227	1310	1397
120	1489	1586	1687	1795	1907	2026	2150	2280	2416	2560
140	2709	2866	3030	3202	3381	3569	3764	3968	4181	4402
160	4633	4874	5124	5384	5655	5937	6229	6533	6848	7175
180	7514	7866	8230	8608	8999	9404	9823	10256	10705	11168
200	11647	12142	12653	—	—	—	—	—	—	—

(Battelli, 1892.)

Temp. . .	220°C.	240°	260°	280°	300°	320°	340°	360°
Vap. Press.	17,380 mm.	25,170	35,760	50,600	67,620	88,340	113,830	141,870

Interpolate logs of vapour pressures as explained above.

BOILING-POINT OF WATER UNDER VARIOUS BAROMETRIC PRESSURES

Hydrogen scale of temps. Pressures in mms. of mercury at 0°C . ; $g = 980.62$ cms. per sec.² (Regnault's measurements ; reduced by Broch, 1881 ; recalculated by Wiebe, 1893.)

Barometric Height.	0	1	2	3	4	5	6	7	8	9
$^{\circ}\text{C}$.										
680 mm.	96.91	96.95	97.00	97.03	97.07	97.11	97.15	97.20	97.24	97.28
690	97.32	.36	.40	.44	.48	.52	.56	.59	.63	.67
700	97.71	.75	.79	.83	.87	.91	.95	.99	98.03	98.07
710	98.11	98.14	98.18	98.22	98.26	98.30	98.34	98.38	.42	.45
720	98.49	.53	.57	.61	.65	.69	.72	.76	.80	.84
730	98.88	.91	.95	.99	99.03	99.07	99.10	99.14	99.18	99.22
740	99.25	99.29	99.33	99.37	.41	.44	.48	.52	.56	.59
750	99.63	.67	.70	.74	.78	.81	.85	.89	.93	.96
760	100.00	100.03	100.07	100.11	100.15	100.18	100.22	100.26	100.29	100.33
770	100.37	.40	.44	.47	.51	.55	.58	.62	.66	.69
780	100.73	.76	.80	.84	.87	.91	.94	.98	101.01	101.05

VAPOUR PRESSURE OF MERCURY

In mms. of mercury at 0°C . Reduced from the observations of Hertz, Ramsay and Young, Callendar and Griffiths, Pfundler, Morley, Gebhardt, Cailletet, Colardeau, Rivière. For interpolation from 15° to 270° .

$$\log p = 15.24431 - 3623.932/\theta - 2.367233 \log \theta \quad \dots \quad (\text{A})$$

From 270° to 450°

$$\log p = 10.04087 - 3271.245/\theta - .7020537 \log \theta$$

$\frac{\delta p}{\delta t}$ at the boiling-point = 13.6 mm. per degree (Laby, *Phil. Mag.*, Nov., 1908).

Temp.	Vap. Press.	Temp.	Vap. Press.	Temp.	Vap. Press.	Temp.	Vap. Press.	Temp.	Vap. Press.
0°C .	mm. .00016*	25°	mm. .00168	60°	mm. .0246	250°	mm. .7583	500°	atmos. 8
5	.00026*	30	.00257	80	.0885	300	248.6	600	22.3
10	.00043*	35	.00387	100	.276	356.7	760	700	50
15	.00069	40	.00574	150	2.83	400	1566	800	102
20	.00109	50	.0122	200	17.81	450	3229	880	162

* Extrapolated by formula A.

VAPOUR PRESSURE OF ETHYL ALCOHOL

Vap. press. in mms. of mercury at 0°C . Calculated by Bunsen from Regnault's results (1862), which are in good agreement with the mean of those of Ramsay and Young (1886), and Schmidt (1891).

Regnault, Vapour press. at -20° , 3.34 mm.; at -10° , 6.47 mm.

Temp.	0	1	2	3	4	5	6	7	8	9
0°C .	12.73	13.65	14.6	15.59	16.62	17.7	18.84	20.04	21.31	22.66
10	24.08	25.59	27.19	28.9	30.7	32.6	34.6	36.8	39.0	41.4
20	44.0	46.7	49.5	52.5	55.7	59.0	62.5	66.2	70.1	74.1
30	78.4	—	—	—	—	—	—	—	—	—

(Ramsay and Young, 1886.)

Temp.	30°C .	40°	50°	60°	70°	80°	100°	120°	140°	160°
Press.	78.1 mm.	133.4	219.8	350.2	541	812	1692	3220	5670	9370

Interpolate logs of vapour pressures as explained on p. 40.

VAPOUR PRESSURES

VAPOUR PRESSURES OF ELEMENTS

p = vapour pressure in mms. of mercury at 0° C. lat. 45° and sea-level ($g = 980.62$) (i.e. 1 mm. Hg = 1333.2 dynes per sq. cm). If followed by *at.*, p is in atmospheres; θ = absolute temp. (A.); t = temp. in $^{\circ}$ C.; (s) solid; (l) liquid. The thermometry is in many cases somewhat dubious.

Interpolate logs of vapour pressures as explained on p. 40.

Argon	t	-121° C.	-128.6	-129.6	-134.4	-135.1	-136.2	-138.3	-139.1	—
(Olszewski, 1895)	p	50.6 at.	38.0	35.8	29.8	29.0	27.3	25.3	23.7	—
Argon	θ	78° 9 A.	86.9	97.9	107.3	155.6	= crit. temp.			
Krypton	θ	110° 5 A.	121.3	135.2	147.3	—	210.5	= crit. temp.		
Xenon	θ	148° 9 A.	163.9	182.9	199.6	—	—	287.8	= crit. temp.	
(Ramsay & Travers)	p	300 mm.	760	2000	4000	40,200	41,240	43,500	—	—
Bromine	t	-16° 6 C.	-12.0	-5.0	8.2	16.9	23.4	40.5	51.9	58.7
(Ramsay & Young, 1886)	p	20 mm.	30	50	100	150	200	400	600	760
Chlorine	t	-80° C.	-60°	-40	-33.6	-20	0	10	20	30
(Knietzsch, 1890)	p	62.5 mm.	210	560	760	1.84 at.	3.66	4.95	6.62	8.75
Iodine (Baxter, Hickey, & Holmes, 1907)	t	0° C.	15	30	55	85	117	137	160.9	185.3
	p	0.2 mm.	.131	.469	3.08	20	100	200	400	760
Hydrogen (Travers & Jaquerod, 1902)	t	-258° 2 C.	-256.7	-255.7	-255.0	-254.3	-253.7	-253.2	-252.9	H. Scale
	p	100 mm.	200	300	400	500	600	700	760	—
Helium	θ	1° 2 A.	4.3	—	—	Neon (Travers & Jaquerod, '02)				
(Onnes, 1911)	p	0.2 mm.	760	—	—	15° 65 A. (s) 20.4 (s) He Scale				
						2.4 mm. 12.8				
Mercury	See p. 41.					Ra. Emanation See p. 103.				
Nitrogen (Baly, 1900	θ	62° 5 A.	67.8	72.4	77.3	80	83	86	89	91
Fischer & Alt., 1902)	p	86 mm.	200	400	760	1013	1386	1880	2465	2916
Oxygen (Jaquerod, Travers, & Senter, 1902)	θ	79° 1 A.	82.1	84.4	86.3	87.9	89.3	90.1	90.6	H. Scale
	p	200 mm.	300	400	500	600	700	760	800	—
Phosphorus	t	165° C.	170	180	200	209	219	226	230	287.3
(Schrötter, 1848)	p	120 mm.	173	204	266	339	359	393	514	760
Sulphur (Ruff & Graff, '08; B., 1899; C., 1899)	t	50° C.	100	147	211	400	444.5	$\delta t/\delta p = 0^{\circ} 09/\text{mm.}$ near B.P. (see p. 50).		
	p	0.003 mm.	.0089	.192	3.14	c. 372	760			

VAPOUR PRESSURES OF COMPOUNDS

For a complete list, see Schenck in L.B.M.

Hydrochloric acid	t	-73° 3 C.	-45.5	-23.3	-3.9	4.0	9.2	13.8	22.0	33.4
(F., 1845; Ansdell, 1880)	p	1.8 at.	6.3	12.8	23.1	29.8	33.9	37.7	45.7	58.8
Sulphuretted hydrogen	t	-25° C.	-15	-5	0	10	30	50	60	70
(R., 1862)	p	4.93 at.	6.84	9.3	10.8	14.3	23.7	36.6	44.4	53.1
Sulphur dioxide	t	-30° C.	-20	-10	0	10	20	30	40	50
(Regnault, 1862)	p	.39 at.	.63	1.00	1.53	2.26	3.24	4.52	6.15	8.19
Ammonia, NH ₃	t	-80° C.	-77.6	-70.4	-64.4	-60.8	-54.4	-46.2	-39.8	-33.0
(Brill, 1906)	p	35.2 mm.	44.1	74.9	116.0	157.6	239.5	403.5	568.2	761
Nitrous oxide, N ₂ O	t	-80° C.	-60	-40	-20	-10	0	10	20	40
(Cailliet, '78; R., '62)	p	1.9 at.	5.05	11.0	23.1	28.9	36.1	44.8	55.3	83.4
Nitric oxide, NO	t	-176.5° C.	-167	-138	-129	-119	-110	-105	-100.9	-97.5
(Olszewski, 1885)	p	.024 at.	.182	.54	10.6	20.0	31.6	41.0	49.9	57.8
Nickel carbonyl, NiCO	t	-9° C.	-7	-2	0	10	16	20	30	—
(D. & Jones, 1903)	p	24.3 mm.	104.3	129.1	144.5	215.0	283.5	329.5	462	—

Interpolate logs of vapour pressures as explained on p. 40.

VAPOUR PRESSURES OF COMPOUNDS (*contd.*)

Interpolate logs of vapour pressures as explained on p. 40.

Carbon dioxide	t	-130° C. (s)	100 (s)	-80 (s)	-65 (s)	-56.4 †	-65 (l)	40 (l)	-20 (l)	-10 (l)
(Zeleny & Smith, 1906) . . .	p	2.5 mm.	119	657	2100	3910	2508	7510	14,830	19,630
Carbon bisulphide	t	-20° C.	-10	0	10	20	40	60	80	100
(Regnault, 1862)	p	47.3 mm.	79.4	128	198	298	618	1164	2033	3325
Chloroform, CHCl ₃	t	20° C.	30	40	50	60	70	80	90	100
(Regnault, 1862)	p	160.5 mm.	248	369	535	755	1042	1408	1865	2429
Carbon tetrachloride, CCl ₄ . .	t	-20° C.	-10	0	10	20	40	60	80	100
(R., 1862)	p	9.8 mm.	18.47	32.9	56	91	215	447	843	1467
Acetylene, C ₂ H ₂	t	-90° C. (s)	-85 (s)	-81	-70	-50	-23.8	0	20.2	36.5
(Villard, 1895)	p	.69 at.	1.00	1.25	2.22	5.3	13.2	26.05	42.8	61.6 (M.)
Benzene, C ₆ H ₆	t	-10° C.	0	10	20	40	60	80	100	120
(Young, 1889)	p	14.8 mm.	26.5	45.4	74.6	181.1	389	754	1344	2238
Aniline, C ₆ H ₅ NH ₂	t	101.9 C.	119.4	138.7	151.5	161.1	168.7	175.0	180.8	183.9
(Kahlbaum, 1898)	p	50 mm.	100	200	300	400	500	600	700	760
Bromnaphthalene	t	215° C.	220	230	240	250	260	270	275	280.4
(C ₁₀ H ₇ Br (Ra. & Y., 1885) . .	p	158.9 mm.	181.8	236.0	303.4	386.4	487.4	608.8	677.9	760
Me. alcohol, CH ₃ OH	t	-10° C.	0	17	20	30	50	80	120	150
(R., '62; Ra. & Y.; Ri., '86) .	p	14.8 mm.	28.5	78.3	88.7	150	381.7	1238	4312	9361
n. propyl alcohol, † C ₃ H ₇ OH .	t	0° C.	10	17	30	40	60	80	100	120
(Ra. & Y.; S.; Ri., '86) . . .	p	3.9 mm.	7.8	12.4	28.2	51.4	157	389	843	1668
Iso-butyl alcohol †	t	10° C.	17	20	40	60	80	100	108	120
(C ₄ H ₉ OH (Ri., '86; S., '91) . .	p	4.1 mm.	6.8	8.1	30.3	94.2	245	569	760	1195
Iso-amyl alcohol †	t	17° C.	30	40	50	60	80	100	120	130
(C ₅ H ₁₁ OH (Ri., '86; S., '91) .	p	1.78 mm.	4.68	9.33	17.4	32.0	151	234	522	741
Formic acid, † CH ₂ O ₂	t	0° C.	10	17	20	30	40	70	80	101
(S., 1891; K., 1898)	p	10.2 mm.	18.4	26.3	31.6	51.3	79.4	266	373	760
Acetic acid, † C ₂ H ₄ O ₂	t	17° C.	30	50	70	90	110	130	150	200
(Ra. & Y.; Ri., '86; S., '91) .	p	9.8 mm.	20.6	56.2	133	288	582	1058	1847	5905
Propionic acid, † C ₃ H ₆ O ₂ . .	t	15° C.	17	20	30	40	60	70	80	140
(Ri., '86; S., '91; K., '98) . .	p	1.7 mm.	2.0	2.45	4.9	9.1	28.2	46.1	74.5	760
Butyric acid, † C ₄ H ₈ O ₂	t	17° C.	20	30	50	70	90	110	130	150
(Ra. & Y., '86; S., '91; K., '94) .	p	.52 mm.*	.66*	1.4	5.2	16.2	44.9	111	245	497
Iso-butyric acid, † C ₄ H ₈ O ₂ . .	t	17° C.	30	50	70	90	110	130	150	153.5
(Ri., '86; S., '91; K., '94) . .	p	.88 mm.*	1.9	8.2	25.1	67.6	162	347	684	760
Methyl formate †	t	-20° C.	-10	0	10	20	40	60	80	100
(CHO ₂ CH ₃ (Y. & T., '93) . .	p	67.7 mm.	117.6	195	309	476	1029	1990	3497	5782
Methyl butyrate †	t	-10° C.	0	10	20	40	60	80	100	—
(C ₄ H ₇ O ₂ CH ₃ (Y. & T., '93) .	p	3.55 mm.	7.3	13.8	24.5	69.2	167.5	361	701	—
Methyl isobutyrate †	t	-10° C.	0	10	20	40	60	80	100	120
(C ₄ H ₇ O ₂ CH ₃ (Y. & T., '93) .	p	6.22 mm.	12.15	22.4	38.9	104.7	244	505	956	1660
Ethyl acetate †	t	-20° C.	-10	0	10	20	40	60	80	100
(C ₂ H ₃ O ₂ C ₂ H ₅ (Y. & T., '93) .	p	6.5 mm.	12.9	24.3	42.7	72.8	186	415	833	1515
Ethyl propionate †	t	-10° C.	0	10	20	40	60	80	100	120
(C ₂ H ₅ O ₂ C ₂ H ₅ (Y. & T., '93) .	p	4.05 mm.	8.3	15.5	27.7	77.9	188.0	403.6	785	1388
Propyl acetate †	t	-10° C.	0	10	20	40	60	80	100	120
(C ₂ H ₅ O ₂ C ₃ H ₇ (Y. & T., '93) .	p	3.6 mm.	7.4	13.9	25.1	70.8	172	373	724	1288
Ethyl ether, (C ₂ H ₅) ₂ O	t	-10° C.	0	10	20	40	60	80	100	193.8 †
(Young, 1910)	p	112.3	184.9	290.8	439.8	921	1734	2974	4855	27,060

Interpolate logs of vapour pressure as explained on p. 40.

* Extrapolated.

† The vapour pressures here given have been graphically interpolated from the observers' values. B., Bodenstein; C., Callendar; D., Dewar; F., Faraday; K., Kahlbaum; M., Mackintosh; R., Regnault; Ra. and Y., Ramsay and Young; Ri., Richardson; S., Schmidt; Y. and T., Young and Thomas.

‡ Triple point.

|| Critical temp.

GAS THERMOMETRY

GAS THERMOMETRY

The standard thermometric scale of the International Committee of Weights and Measures (1887) is that of the constant-volume hydrogen thermometer, the hydrogen being taken at an initial pressure at 0° C. of 1000 mms. of mercury measured at 0° C. sea-level and lat. 45° (= 1'3158 standard atmosphere).

THERMODYNAMIC TEMPERATURE OF THE ICE-POINT

Method.	H ₂	N ₂	Air.	CO ₂	Computer.
	°	°		°	
From Joule-Thomson effect	273'14	273'09	—	273'05	Callendar, 1903
Extrapolation to zero pressure	273'07	273'09	—	—	Berthelot and
(see p. 54)			°		Chappuis, 1907
From Joule-Thomson effect	273'05	(273'17)	273'19	273'10	Berthelot, 1907
" " "	273'06	273'25	273'27	273'12	Buckingham, 1908
" " "	273'13	273'14	—	—	Rose-Innes, 1908

General mean = 273°·13.

THERMODYNAMIC CORRECTIONS TO GAS SCALES OF TEMPERATURE

The corrections to both the constant-pressure (C.P.) and the constant-volume (C.V.) scales are either (1) derived from characteristic equations of state (Callendar, 1903; Berthelot, 1907), or (2) in the case of the C.P. thermometer, computed from the Joule-Thomson effect; whence from these C.P. corrections and a knowledge of the compressibility of the gas under different conditions the C.V. corrections can be calculated. Chappuis (1907)* has experimentally compared the C.P. and C.V. H. and N. thermometers each with mercury thermometers. The values below are based on computations by Callendar (*Phil. Mag.*, 1903), Berthelot* (from Chappuis' data 1907), Onnes and Braak (1907 and 1908), Rose-Innes (*Phil. Mag.*, 1908), and Buckingham (1908).† There is some divergence among the different computations for hydrogen; the agreement is much better in the case of nitrogen. The thermodynamic correction to the C.V.H. thermometer is negligible, and with nitrogen also at extreme temps. the correction is less than the error of working in modern gas thermometry. The values for air are a little smaller than for nitrogen; for helium they are slightly larger than for hydrogen except at the lowest temperatures, when the helium corrections are the smaller. New experiments on the Joule-Thomson effect are needed. ‡ (+) means that the correction has to be added to the gas scale temperature to give the thermodynamic temperature. The correction is proportional to the initial pressure of the gas in the thermometer.

* *Trav. et Mém. Bureau Intl.* 1907.

† *Bull. Bureau of Standards.* 1908.

‡ See Dalton, *Proc. Konink. Akad. Weten. Amsterdam*, April, 1909.

t° C.	Const. Pressure P = 1000 mm.		Const. Volume P at 0° = 1000 mm.		t° C.	Const. Pressure P = 1000 mm.		Const. Volume P at 0° = 1000 mm.	
	H ₂	N ₂	H ₂	N ₂		H ₂	N ₂	H ₂	N ₂
-240°	+1°·2 (?)	—	+°·18	—	70°	-°·003	-°·019	-°·001	-°·004
-200	+°·26	—	+°·06	—	80	-°·002	-°·014	-°·000	-°·003
-150	+°·10	+1°·3	+°·033	+°·26 (?)	90	-°·001	-°·007	-°·000	-°·002
-100	+°·04	+°·40	+°·010	+°·10 (?)	100	0	0	0	0
-50	+°·02	+°·12	+°·005	+°·03	200	+°·014	+°·12	+°·004	+°·04
0	0	0	0	0	300	+°·034	+°·28	+°·011	+°·10
10	-°·001	-°·009	-°·000	-°·002	400	+°·07 (?)	+°·46	+°·018 (?)	+°·17
20	-°·002	-°·017	-°·000	-°·004	450	+°·09 (?)	+°·56	+°·02 (?)	+°·19
30	-°·003	-°·021	-°·001	-°·005	600	—	+°·87	—	+°·3
40	-°·003	-°·023	-°·001	-°·006	800	—	+1°·3	—	+°·5
50	-°·003	-°·024	-°·001	-°·007	1000	—	+1°·8	—	+°·7
60	-°·003	-°·022	-°·001	-°·006	1200	—	+2°·3	—	+1°·0

MERCURY THERMOMETRY

CORRECTIONS TO REDUCE MERCURY-IN-GLASS SCALE TEMPS. TO GAS SCALE TEMPS.

The values for the English Kew glass (which is a lead potash silicate) are due to Harker (1906); the verre dur corrections are given by the International Bureau; those for the Jena glasses by Grützmacher. The method at Kew is to determine the ice-point correction before an observation is made. The other glasses have their ice-point or zero depressions determined immediately after each temperature reading. See Guillaume's "Thermométrie de Précision," Paris, 1889, and Chree's "Notes on Thermometry," *Phil. Mag.*, 1898. The French glass, verre dur, is used by Tonnelot of Paris. The normal glass, Jena 16^{'''}, may be known by the presence of a thin violet line near the surface. Jena 59^{'''} is a borosilicate (p. 74).

Temp.	Kew Glass.	Verre Dur.	Jena 16 ^{'''} .	Jena 59 ^{'''} .	Temp.	Verre Dur.	Jena 16 ^{'''} .	Jena 59 ^{'''} .
	$t_H - t_{K.G.}$	$t_H - t_{V.D.}$	$t_H - t_{16''}$	$t_H - t_{59''}$		$t_N - t_{V.D.}$	$t_N - t_{16''}$	$t_N - t_{59''}$
-20°	—	+°17	+°19	+°10	110°	+°04	+°03	-°00
0	0°	0	0	0	120	+°06	+°05	-°02
10	-°00	-°05	-°06	-°02	130	+°07	+°07	-°04
20	-°00	-°03	-°09	-°04	140	+°07	+°09	-°08
30	+°005	-°10	-°11	-°04	150	+°06	+°10	-°13
40	+°01	-°11	-°12	-°04	160	+°03	+°10	-°19
50	+°01	-°10	-°11	-°03	170	0	+°08	-°28
60	+°01	-°09	-°10	-°02	180	-°04	+°06	-°39
70	+°015	-°07	-°08	-°01	190	-°09	+°02	-°52
80	+°02	-°05	-°06	-°00	200	-°13	-°04	-°67
90	+°025	-°03	-°03	-°00	250	—	-°63	-1°7
100	0	0	0	0	300	—	-1°91	-4°1

DEPRESSION OF ZERO OF MERCURY THERMOMETERS

The values indicate the zero depressions after the thermometer has been heated to the temp. stated. They have been determined by Guillaume, Thiesen, Schloesser, and Böttcher because of the impossibility in practice of interrupting a series of temperature measurements to take a number of zero readings (see above).

Temp.	Verre Dur.	Jena 16 ^{'''} .	Jena 59 ^{'''} .	Temp.	Verre Dur.	Jena 16 ^{'''} .	Jena 59 ^{'''} .
10° C.	°008	°005	°005	60° C.	°060	°039	°024
20	°017	°011	°009	70	°071	°048	°027
30	°027	°017	°014	80	°084	°057	°030
40	°037	°024	°017	90	°097	°066	°033
50	°048	°031	°021	100	°111	°077	°035

STEM-EXPOSURE OR EMERGENT-COLUMN CORRECTION

The table below gives the (additive) "stem-exposure" correction for (1) the ordinary solid-stem thermometer, and (2) the German pattern sleeve-thermometer, which has a fine capillary in an outer glass tube. Both thermometers are of Jena 16^{'''} glass, with degree intervals about 1 mm. long.

t is the indicated temperature, and t_{aux} the temperature of an auxiliary thermometer whose bulb is 10 cms. from and on a level with the mid-point of the exposed stem. The auxiliary thermometer must be shielded from the source of heat. (See Watson's "Practical Physics," and Rimbach, *Zeit. f. Inst.*, 10, 1890.)

No. of degree divs. of exposed thread.	Solid Stem; Scale on Stem.						Sleeve Thermometer; Enclosed Scale.						No. of degree divs. of exposed thread.
	$t - t_{aux}$												
	70° C.	80°	100°	120°	140°	180°	70° C.	80°	100°	120°	140°	180°	
10	°02	°03	°07	°11	°17	°27	°01	°01	°04	°07	°10	°17	10
20	°13	°15	°22	°29	°38	°53	°08	°12	°19	°25	°28	°40	20
30	°24	°28	°39	°48	°59	°78	°25	°28	°36	°42	°48	°66	30
40	°35	°41	°56	°68	°82	1°04	°30	°35	°48	°60	°67	°92	40
60	°57	°66	°89	1°09	1°25	1°58	°52	°60	°79	°99	1°11	1°46	60
80	°80	°91	1°21	1°52	1°71	2°15	°75	°87	1°15	1°38	1°53	1°98	80
100	1°02	1°18	1°56	1°97	2°18	2°70	°98	1°12	1°47	1°82	2°03	2°55	100
120	—	—	1°98	2°43	2°69	3°26	—	—	1°88	2°28	2°49	3°13	120

ELECTRICAL THERMOMETRY

PLATINUM THERMOMETRY

TO REDUCE PT-SCALE TEMPS. (t_{pt}) TO CONST. VOL. N-SCALE TEMPS. (t)

Callendar's "difference formula" for the difference between the nitrogen-scale temp. (t) and the Pt-scale temp. (t_{pt}) is $t - t_{pt} = \delta \cdot t(t - 100)10^{-4}$, where δ is close to 1.5. Pt-scale temps. result from assuming a linear relation $R_{pt} = R_0(1 + \alpha t_{pt})$ between temp. and the electrical resistance (R) of Pt; α is the mean coefficient for the range 0° to 100° . The "difference formula" gives the correction yielded by the truer parabolic relation $R_t = R_0(1 + \alpha t + \beta t^2)$. Pt thermometers should not be used above 1200° C. (See Callendar, *Phil. Mag.*, 1899, 1, p. 191; 2, p. 519, Camb. Sci. Inst. Co.'s list "Technical Thermometry"; and (for bibliography), Waidner and Burgess, *Bull. Bur. of Standards*, 1909.)

$$\delta = 1.50.$$

(Harker, *Phil. Trans.*, 1904.)

Pt Temps. t_{pt}	0	20	40	60	80	100	120	140	160	180
	t	t	t	t	t	t	t	t	t	t
-200°	—	-172°·9	-154°·1	-135°·2	-116°·2	-97°·13	-77°·92	-58°·61	-39°·18	-19°·65
0	0°	19°·76	39°·64	59°·64	79°·76	100	120°·4	140°·9	161°·5	182°·3
+200	203°·1	224°·2	245°·4	266°·7	288°·1	309°·8	331°·5	353°·4	375°·5	397°·8
400	420°·2	442°·8	465°·5	488°·5	511°·6	534°·9	558°·4	582°·1	606°·0	630°·1
600	654°·4	679°·0	703°·7	728°·7	754°·0	779°·4	805°·2	831°·2	857°·4	884°·0
800	910°·8	937°·9	965°·3	993°·0	1021	1050	1078	1107	1137	1167
1000	1197	1228	1259	1290	1323	1355	—	—	—	—

TO CALCULATE THE CHANGE Δt IN THE N-SCALE TEMP. (t) FOR A CHANGE OF ± 0.1 IN δ

t	Δt	t	Δt	t	Δt	t	Δt	t	Δt	t	Δt
-200°	°·060	-60°	°·010	80°	°·002	250°	°·038	600°	°·30	950°	°·8
-180	°·050	-40	°·006	100	0	300	°·060	650	°·36	1000	°·9
-160	°·042	-20	°·002	120	°·002	350	°·088	700	°·42	1050	°·10
-140	°·034	0	0	140	°·006	400	°·120	750	°·49	1100	°·11
-120	°·026	20	°·002	160	°·010	450	°·158	800	°·56	1150	°·12
-100	°·020	40	°·002	180	°·014	500	°·20	850	°·64	1200	°·13
-80	°·014	60	°·002	200	°·020	550	°·25	900	°·72	1250	°·14

HIGH TEMPERATURES

(See Burgess and Le Chatelier's "High Temperature Measurements, 1912.")

For the measurement of high temperatures (say above 1200° C., which is about the present upper experimental limit of the gas scale) the instruments in general use are thermo-junctions and optical or radiation pyrometers. Both involve extrapolation. Thermo-couples have been used up to the temperature of the melting-point of platinum ($c. 1750^\circ$). At high temperatures thermo-junctions yield rather lower results than do optical pyrometers, e.g. see the M.P.'s of Pd and Pt on p. 49.

THERMO-ELECTRIC THERMOMETRY

Temperature readings with thermo-couples are reduced by one of the formulae: (a) $E = a + bt + ct^2$, (b) $E = mt^n$, or $\log E = n \log t + m'$, E being the e.m.f. generated, and t the temperature of the hot junction, the cold junction being at 0° . Up to about 1200° these formulae with suitable constants agree to within 2° for the usual 10% (Pt, Pt - Rh) and (Pt, Pt - Ir) couples, but above 1200° formula (b) yields the higher results, e.g. see the melting-points of Pd and Pt on p. 49. The thermo-e.m.f.'s of these Pt couples gradually diminish with prolonged heating. The values of the constants below are only average values.

E IN MICRO VOLTS (10^{-6} VOLT)

	Couple.	a	b	c	n	m'
Cold junction at 0° C.	Pt and (90 Pt, 10 Rh)	-307*	8.1*	°·0017*	1.19	°·52
	Pt and (90 Pt, 10 Ir)	-550*	14.8*	°·0016*	1.10	°·89
	Cu and Constantan †	—	—	—	1.14	1.34
	Cu and Fe	0	10.34	-°·0183	—	—

* These constants are not suitable for temperatures below 300° . † Eureka, 60 Cu, 40 Ni.

THERMO-ELECTRIC THERMOMETRY (contd.)

The following are the readings in 10^{-5} volt determined at the National Physical Laboratory for a Pt-Rh and a Pt-Ir couple, each having the cold junction at 0°C . The values only hold for the particular couples.

Couple.	Temp.	0	50	100	150	200	250	300	350	400	450
Pt and (90 Pt, 10 Rh)	0°C .	0	23	51	83	119	158	199	242	286	331
	500	377	423	470	518	567	617	668	720	773	826
	1000	880	935	991	1048	1106	1165	1225	1286	1348	—
Pt and (90 Pt, 10 Ir)	0	0	58	125	195	268	343	420	498	577	657
	500	737	818	899	981	1064	1147	1231	1315	1400	1485
	1000	1571	1657	1744	1831	1919	2007	2096	2185	2275	—

THERMO-E.M.F.'S AGAINST PLATINUM IN MICRO VOLTS (10^{-6} VOLT)

One junction at 0°C . The current flows across the other junction from the metal with the (algebraically) smaller value to the other metal. (See Watson's "Physics" and Henning in L.B.M.)

Metal.	-190°	$+100^{\circ}$	Metal.	-190°	$+100^{\circ}$	Metal.	-190°	$+100^{\circ}$
Aluminium	+ 390	+ 380	Lead.	+ 210	+ 410	Tantalum	—	+ 330
Antimony.	—	+ 4700	Magne-	—	—	Tin . . .	+ 200	+ 410
Bismuth .	+ 12300	- 6500	sium .	+ 330	+ 410	Zinc . . .	- 120	+ 750
Cadmium .	+ 60	+ 900	Mercury	—	0	Brass . . .	—	+ 400
Cobalt . .	—	- 1520	Nickel .	+ 2220	- 1640	Constantan*	—	- 3140
Copper . .	- 200	+ 740	Pallad-	—	—	German sil-	—	—
Gold . . .	- 120	+ 730	dium .	+ 790	- 560	ver†. . .	—	+ 1000
Iron . . .	- 2900	+ 1600	Silver .	- 140	+ 710	Manganin‡	—	+ 570

* Eureka, 60 Cu, 40 Ni.

† 60 Cu, 15 Ni, 25 Zn.

‡ 84 Cu, 4 Ni, 12 Mn.

RADIATION AND OPTICAL THERMOMETRY

Most radiation thermometers depend upon either (1) the Stefan-Boltzmann law, $E = K(\theta^4 - \theta_0^4)$, where E is the total energy (all wave-lengths) radiated per sec. by a black body at absolute temp. θ to surroundings at absolute temp. θ_0 , and K is a const. ($K = 5.7 \times 10^{-12}$ watts per cm^2 per 1° —see p. 65); or (2) Wien's equation connecting the temperature with the intensity of some particular wave-length of light

emitted (p. 65). The Wien equation is, Intensity $I = c_1 \lambda^{-5} e^{-\frac{c_2}{\lambda T}}$, where λ is the wave-length, T is the "black body" temp. on the absolute scale, c_1 and c_2 are constants, and e is the base of the Napierian logarithms. Both equations give results which agree very accurately with the gas scale over the calibrated range 0° to 1200°C . Up to about 1500° radiation thermometers are, in practice, almost always graduated empirically, usually against a thermo-couple.

The "black body" temperature of a radiating substance is the temperature at which an ideal black body would emit radiation of the same intensity as that from the substance, the radiation considered being of some particular wave-length. A perfectly black body absorbs all the radiation which falls upon it; it is destitute of reflecting power. Coal, carbon, metals which when heated tarnish with a black oxide, enclosed furnaces and muffles at a uniform temperature, all conform very nearly to this definition. When a pyrometer is sighted upon a body which is not "black," the temperature recorded—the "black body" temperature—will be lower than the true temperature to an extent which increases with the reflecting power of the body, e.g. if platinum and carbon have equal "black body" temperatures, their actual temperatures may differ by 180° or so at 1500° .

TEMPERATURE AND COLOUR OF FIRE

Appearance .	Red—just visible.	Dull Red.	Cherry Red.	Orange.	White.	Dazzling White.
Temperature.	c. 500°C .	c. 700°	c. 900°	c. 1100°	c. 1300°	c. 1500°

For standard temperatures for thermometer calibration, see p. 50.

MELTING AND BOILING POINTS

MELTING AND BOILING POINTS OF THE ELEMENTS

For an account of temperature measurements, see p. 46. For melting and boiling points of chemical compounds, see p. 109; of fats and waxes, see p. 50.

Element.	Melting Point.	Observer.	Boiling Point at 760 mms.	Observer.
Aluminium . . .	657° C.	Holborn and Day, 1900	1800° C.	Greenwood, 1909
Antimony . . .	630	" "	1440	Greenwood, 1909
Argon . . .	-188	Ramsay and Travers, 1901	-186	—
Arsenic . . .	volatilizes	—	{sublimes}	—
Barium . . .	850	Guntz, 1903	450	—
Beryllium . . .	c. 1430	Just and Mayer, 1909	—	—
Bismuth . . .	269	Callendar, 1899	1420	Greenwood, 1909
Boron . . .	2000 to 2500	Weintraub, 1909	{sublimes}	—
Bromine . . .	-7.3	van der Plaats, 1886	3500 (?)	van der Plaats, 1886
Cadmium . . .	321	Holborn and Day, 1900	63	D. Berthelot, 1902
Cæsium . . .	26.4	Eckardt and Graefe, 1900	778	Ruff & Johannsen, 1906
Calcium . . .	780	Ruff and Plato, 1903	670	—
Carbon . . .	4000 (?)	(Calculated) McCrac, 1906	—	—
Cerium . . .	623	Muthmann & Weiss, 1904	—	—
Chlorine . . .	-102	Olzewski	-33.6	Regnault, 1863
Chromium . . .	1520	Bureau of Standards	2200	Greenwood, 1909
Cobalt . . .	1480	Bureau of Standards	—	—
Copper . . .	{ 1084 *	Holborn and Day, 1900	2310	Greenwood, 1909
Erbium . . .	{ 1083	Day and Sosman, 1910	—	—
Fluorine . . .	-223	Moissan and Dewar, 1903	—	Moissan & Dewar, 1903
Gallium . . .	30.2	L. de Boisbaudran, 1876	-187	—
Germanium . . .	960	Biltz, 1911	—	—
Gold . . .	{ 1063	Holborn and Day, 1901	2530 (?)	—
Helium . . .	{ 1062 †	Day and Sosman, 1910	—	—
Hydrogen . . .	below -272	Onnes, 1911	-268.8	Onnes, 1911
Indium . . .	-259	Travers, 1902	-252.7	Travers, 1902
Iodine . . .	155	Thiel, 1904	1000 (?)	—
Iridium . . .	113	Lean & Whatmough, 1898	184.4	Drugmann & Ramsay, 'co
Iron . . .	2290	Mendenhall & Ingersoll, '07	2550 (?)	—
Krypton . . .	1530	Bureau of Standards	2450	Greenwood, 1909
Lanthanum . . .	-169	Ramsay, 1903	-151.7	Ramsay, 1903
Lead . . .	810	Muthmann & Weiss, 1904	—	—
Lithium . . .	327	Holborn and Day, 1900	1525	Greenwood, 1909
Magnesium . . .	186	Kahlbaum, 1900	>1400	Ruff & Johannsen, 1906
Manganese . . .	633	Heycock and Neville, 1895	1120	Greenwood, 1909
Mercury . . .	1260	Bureau of Standards	1900	Greenwood, 1909
Molybdenum . . .	-38.80	Chappuis, 1900	356.7	Callendar, 1899
Neodymium . . .	2450	Pirani & Meyer, 1912	3200 (?)	—
Neon . . .	840	Muthmann & Weiss, 1904	—	—
Nickel . . .	—	—	-239	Dewar, 1901
Niobium . . .	1452 †	Day and Sosman, 1910	2330 (?)	—
Nitrogen . . .	1950	von Bolton, 1907	—	—
	-210.5	Fischer and Alt, 1903	-195.7	Fischer & Alt, 1903

* In reducing atmosphere; 1062° in air.

† Const. vol. N. thermometer.

MELTING AND BOILING POINTS OF THE ELEMENTS (*contd.*)

Element.	Melting Point.	Observer.	Boiling Point at 760 mms.	Observer.
Osmium . . .	2700° C.	—	—	—
Oxygen . . .	-219	Dewar, 1911	-182°·9 C.	Travers, 1902
Palladium*—	1549 ‡	Day and Sosman, 1910	2540	—
thermo-jn. (a)	1535	Holborn & Henning, 1905	—	—
optical therm.	1549	—	—	—
" . . .	1545	Nernst & Wartenberg, 1906	—	—
" . . .	1582	Holborn & Valentiner, 1907	—	—
thermo-jn. (a)	1530	Waidner & Burgess, 1907	—	—
" (b)	1543	" "	—	—
optical therm.	1546	" "	—	—
Phosphorus . .	44·1 ₇₆₀	Hulett, 1899	287	Schrötter, 1848
Platinum*—	—	—	—	—
thermo-jn. (a)	1710	Harker, 1905	2450 (?)	—
" (a)	1710	Holborn & Henning, 1905	—	—
optical therm.	1729	" "	—	—
" . . .	1750	Nernst & Wartenberg, 1906	—	—
" . . .	1789	Holborn & Valentiner, 1907	—	—
thermo-jn. (a)	1706	Waidner & Burgess, 1907	—	—
" (b)	1731	" "	—	—
optical therm.	1770	" " 1909	—	—
Potassium . . .	62·5	Holt and Sims, 1894	758	Ruff & Johannsen, 1905
Praseodymium .	940	Muthmann and Weiss, 1904	—	—
Radium . . .	700	Curie and Debierne, 1910	—	—
Rhodium . . .	1907	Mendenhall & Ingersoll, '07	2500 (?)	—
Rubidium . . .	38·5	Erdmann and Köthner, 1896	696	Ruff & Johannsen, 1905
Ruthenium . . .	1900 (?)	—	2520 (?)	—
Samarium . . .	1350	—	—	—
Selenium . . .	217	Saunders, 1900	690	Berthelot, 1902
Silicon . . .	1200 (?)	—	3500 (?)	—
Silver . . .	{ 962 † 960 †	{ Holborn and Day, 1900 Day and Sosman, 1910 }	{ 1955 877 742 }	{ Greenwood, 1909 Ruff & Johannsen, 1905 Permann, 1889 }
Sodium . . .	97·0	Kurnakow & Puschin, 1902	—	—
Strontium . . .	900	—	—	—
Sulphur . . .	{ 115 rhombic 119 monoclinic }	{ — — — }	{ 444·55 (c.p. air) 444·7 (c.v. N) 444·53 (c.p. N) }	{ Eumorfopoulos, 1908 (corrected, 1909) Chappuis & Harker, 1902 Callendar, 1899 }
Tantalum . . .	2910	Burgess, 1907	—	—
Tellurium . . .	450	Matthey, 1901	1390	Deville and Troost, 1880
Thallium . . .	301	Kurnakow & Puschin, 1901	1280 (?)	Wartenberg, 1907
Thorium . . .	1690	Wartenberg, 1909	—	—
Tin . . .	232	Heycock & Neville, 1895	2270	Greenwood, 1909
Titanium . . .	1800	—	—	—
Tungsten . . .	3500	Gen. Elect. Co. Lab.	3700 (?)	—
Vanadium . . .	1720	—	—	—
Xenon . . .	-140	Ramsay, 1903	-109	Ramsay, 1903
Zinc . . .	418 †	Day and Sosman, 1910	918	Berthelot, 1902
Zirconium . . .	c. 2300	—	—	—

* See section on thermo-electric thermometers, p. 46, for meaning of (a) and (b).

† In reducing atmosphere; 995° in air.

‡ Const. vol. N. thermometer.

STANDARD TEMPERATURES

STANDARD TEMPERATURES

Melting and boiling points of elements will be found on p. 48; of chemical compounds, on p. 109.

B.P. = boiling point at 760 mm.; M.P. = melting point; T.P. = transition point.

Substance.		Temp.	Substance.		Temp.
		° C.			° C.
Hydrogen	B.P.	-253	Zinc *	M.P.	419·4
Oxygen	B.P.	-183	Sulphur *	B.P.	441·7
Carbon dioxide	B.P.	-78·2	Aluminium	M.P.	657
Mercury	M.P.	-38·8	NaCl (Harker)	M.P.	801
Water	M.P.	0	K ₂ SO ₄	M.P.	1070
Na ₂ SO ₄ · 10H ₂ O	T.P.	32·383	Palladium (p. 49)	M.P.	1550
Water	B.P.	100	Platinum (p. 49)	M.P.	1750
Naphthalene *	B.P.	218·0	Tin (Greenwood)	B.P.	2270
Tin *	M.P.	231·9	Arc † (W. & B.) †	—	3700 abs.
Benzophenone *	B.P.	306·0	Arc † (Harker, '08) †	—	3620 abs.
Cadmium *	M.P.	321·0	Sun † (p. 66)	—	5800 abs.

* Const. vol. N. scale, Waidner & Burgess, 1911; W. & B., Waidner & Burgess, 1904.

† Black body temperature.

‡ Positive crater.

EFFECT OF PRESSURE ON BOILING POINTS

$\delta p / \delta t$ is given as mm. Hg per degree C. for pressures not very far removed from 760 mm.

The boiling point in absolute degrees C. of a substance under 760 mm. = $t + c(760 - p)(t + 273)$, where c is a constant for the substance, and t is the B.P. in degrees C. at the pressure p mm. The constant c is the same for chemically similar substances.

(See Young, "Fractional Distillation.")

Substance.	$\delta p / \delta t$	c	Substance.	$\delta p / \delta t$	c	Substance.	$\delta p / \delta t$	c
		$\times 10^{-6}$			$\times 10^{-6}$			$\times 10^{-6}$
Hydrogen	200	—	CCl ₄	23	123	Benzene	23·5	121
Oxygen	77	146	Pentane, n.	25·8	125	Toluene	21·7	120
Carbon dioxide	55	—	Alcohol, methyl	29·6	100	Aniline	19·6	112
Water	27·2	99	" ethyl	30·3	94	Naphthalene	17·1	119
Mercury	13·6	118	" amyl	25	98	Benzophenone	15·8	109
Sulphur *	11·0	114	Ether, ethyl	26·9	121	Acetone	26·4	115

* $t_p = t_{760} + \cdot 0904(p - 760) - \cdot 0452(p - 760)^2$, Harker & Sexton, 1908.

MELTING, FREEZING, AND BOILING POINTS OF FATS AND WAXES

At 760 mm. pressure.

(See Lewkowitsch's treatise.)

Substance.	M.P.	F.P.	Substance.	M.P.	F.P.	Substance.	M.P.	B.P.
	° C.	° C.		° C.	° C.		° C.	° C.
Butter	28-33	20-23	Beeswax	61-64	60-63	Paraffin wax,		
Lard	36-40	27-30	Spermaceti	42-49	42-47	Soft	38-52	350-390
Tallow, beef	40-45	27-35	Stearin	71·6	70	Hard	52-56	390-430
" mutton	44-45	36-41	Naphthalene	80·0	—	Olive oil	—	c. 300

THERMAL CONDUCTIVITIES

The thermal conductivity, k , is given below as the number of (gram) calories conducted per sq. cm. per sec. across a slab of the substance 1 cm. thick, having a temp.-gradient of 1°C. per cm., *i.e.* calorie $\text{cm.}^{-1} \text{sec.}^{-1} \text{temp.}^{-1}$. (See Callendar, "Conduction of Heat," *Encyc. Brit.*, and Winkelmann's "Handbuch der Physik," III, 1906.)

METALS AND ALLOYS

k for most pure metals decreases with rise of temperature; the reverse appears to be true for alloys. If κ be the electrical conductivity and θ the absolute temp., then $k/(\kappa\theta)$ is very approximately a constant for pure metals. (See J. J. Thomson, "Corpuscular Theory of Matter," and Lees, *Phil. Trans.*, 1908.) The electrical conductivity of the same specimen of many of the substances below will be found on p. 81.

Substance.	Temp.	Cond. k.	Observer.	Substance.	Temp.	Cond. k.	Observer.
	$^{\circ}\text{C.}$				$^{\circ}\text{C.}$		
Metals—				Mercury . . .	0	'0148	H. F.
Aluminium* . .	-160	'514	Lees,	" . . .	50	'0189	Weber, '79
" . . .	18	'504	<i>P. T.</i> , '08	" . . .	50	'0177	A., 1864
" . . .	18	'480	J. & D.,	" . . .	17	'0197	R. W., '02
" . . .	100	'492	1900	Nickel . . .	-160	'129	Lees, '08
Antimony . . .	0	'044	Lorenz,	" {97%}	18	'142	J. & D.,
" . . .	100	'040	1881	" {Ni}	100	'138	1900
Bismuth . . .	-186	'025	M., 1907	Palladium . .	18	'168	J. & D.,
" . . .	18	'0194	J. & D.,	" . . .	100	'182	1900
" . . .	100	'0161	1900	Platinum . . .	18	'166	J. & D.,
Cadmium, pure .	-160	'239	Lees, '08	" . . .	100	'173	1900
" . . .	18	'222	J. & D.,	Silver, pure . .	-160	'998	Lees,
" . . .	100	'216	1900	" . . .	18	'974	1908
Copper, pure . .	-160	'079	Lees, '08	" . . .	18	'006	J. & D.,
" . . .	18	'918	J. & D.,	" . . .	100	'992	1900
" . . .	100	'908	1900	Tin, pure . . .	-160	'192	Lees, '08
Gold . . .	18	'700	J. & D.,	" . . .	18	'155	J. & D.,
" . . .	100	'703	1900	" . . .	100	'145	1900
Iron, pure . . .	18	'161	J. & D.,	Tungsten . . .	18	'35	Coolidge
" . . .	100	'151	1900	Zinc, pure . . .	-160	'278	Lees, '08
" wrought . .	-160	'152	Lees, '08	" . . .	18	'265	J. & D.,
" " † . . .	18	'144	J. & D.,	" . . .	100	'262	1900
" " † . . .	100	'143	1900				
" cast ‡ . . .	54	'114	Callendar	Alloys—			
" " ‡ . . .	102	'111		Brass . . .	-160	'181	Lees,
" " § . . .	30	'149	Hall	" . . .	17	'260	1908
" steel {1%}	-160	'113	Lees,	" . . .	18	'054	J. & D.,
" " {C.}	18	'115	1908	Constantan }	100	'064	1900
" " " . . .	18	'108	J. & D.,	(Eureka)¶ }	0	'070	Lorenz,
" " " . . .	100	'107	1900	German silver	100	'089	1881
Lead, pure . .	-160	'092	Lees, '08	" . . .	-160	'035	Lees, '08
" . . .	18	'083	J. & D.,	" . . .	18	'053	J. & D.,
" . . .	100	'082	1900	" . . .	100	'063	1900
Magnesium . .	0 to	'376	Lorenz,	Platinoid . . .	18	'060	Lees, '08
" . . .	100		1881				

* 99% Al. † 1% C., 2% Si, 1% Mn. ‡ 2% C., 3% Si, 1% Mn.

§ 3.5% C, 1.4% Si, .5% Mn. || 70 Cu, 30 Zn. ¶ 60 Cu, 40 Ni.

** 84 Cu, 4 Ni, 12 Mn.

A., Ångström; J. & D., Jaeger & Diesselhorst; M., Macchia; R. W., R. Weber; *P. T.*, *Phil. Trans.*

The values below are mostly at ordinary temperatures. They must be regarded as rough average values in the case of indifferent conductors. Nearly all liquids have very approximately the same conductivity, which in most cases appears to increase with temperature.

* Perp. to cleavage plane. † Average for igneous and sedimentary rocks; see Brit. Ass. Reports. L., Lees, 1892 & 1898; M. & C., Milner & Chattock, 1898; R. W., R. Weber.

In the case of a gas the thermal conductivity $k = 1.603\eta c_v$, where η is the viscosity, and c_v the specific heat at constant volume. Stefan, and Kundt and Warburg have found, in agreement with this formula, that k for air, hydrogen, etc., is constant between the pressures 76 cm. and 1 cm. k increases with the temperature. (See Meyer's "Kinetic Theory of Gases.")

* Mean of five observers. E., Eckerlein, 1900; G., Graetz, 1885; S., Schwarze, 1903; Sc., Schleiermacher, 1889; W., Winkelmann, 1875.

To represent accurately over any considerable range the variation of length (l) with temperature (t) requires for almost all solid substances a parabolic or cubic equation in t . But if the temperature interval is not large, a linear equation $l_t = l_0(1 + \alpha t)$ may be employed; and this gives a definition of the mean coefficient of linear expansion (α) over that temperature range. The coefficient of **cubical expansion** = 3α .

There is little point in tabulating coefficients of higher-powered terms of t , since for a given specimen it is as a rule impossible without measurement to assume with any accuracy anything more definite than the average value of even the first power coefficient (a). Except in a few cases the linear coefficient as defined above increases with the temperature. The values of a subjoined are per degree C., and except when some temperature is specified, for a range round and about 20° C. Some substances expand irregularly, and extrapolation of a may therefore be dangerous. Interpolation of a from the constituent metals must be employed with caution in the case of alloys. (See Winkelmann's "Handbuch der Physik," iii. 1906.)

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COEFFICIENTS OF EXPANSION

COEFFICIENTS OF LINEAR EXPANSION OF SOLIDS (contd.)

Element.	α .	Obs.	Element.	α .	Obs.	Element.	α .	Obs.
	$\times 10^{-6}$			$\times 10^{-6}$			$\times 10^{-6}$	
Aluminium	25.5	V. '93	Copper	16.7	V. '93	Palladium	11.7	S. '03
Antimony	12	F. '69	Gold	13.9	V. '93	Platinum	8.9	B. '88
Bismuth	15.7	V. '93	Iridium	6.5	B. '88	Potassium	83	H. '82
C. (diamond)	1.2	F. '69	Iron (cast)	10.2	D. '02	Selenium, 40°	36.8	F. '69
„ (gas carbon)	5.4	F. '69	„ (wrought)	11.9	H.D. '00	Silver	18.8	V. '93
„ (graphite)	7.9	F. '69	Steel, 10.5 to 11.6	11.6	N.P.L.	Sulphur	6.70	—
Cadmium	28.8	M. '66	Lead	27.6	M. '66	Thallium, 40°	30.2	F. '69
Cobalt	12.3	T. '99	Magnesium	25.4	V. '93	Tin	21.4	M. '66
			Nickel	12.8	T. '99	Zinc, 25.8 to 26.3	26.3	N.P.L.

Substance.	α .	Obs.	Substance.	α .	Obs.
	$\times 10^{-6}$			$\times 10^{-6}$	
Alloys—			Miscellaneous (contd.)		
Aluminium bronze	17.0	N.P.L.	Glass, flint, 45 SiO ₂	7.8	Sc.
Brass (ordy.) c. 66 Cu, 34 Zn	18.9	N.P.L.	„ 8 K ₂ O, 46 PbO	7.8	T.S.S.
Bronze, 32 Cu, 2 Zn, 5 Sn §	17.7	B. '88	„ Jena, 16 ^{III} (see p. 74)	5.7	'96
Constantan (Eureka), 60 Cu, 40 Ni	17.0	N.P.L.	„ 59 ^{III} (see p. 74)	7.2	C. '07
German silver, 60 Cu, 15 Ni, 25 Zn, 50°	18.4	Pf. '72	„ Verre dur (see p. 74)	8.3	—
Gunmetal (Admiralty)	18.1	N.P.L.	Granite	19.8	Ru. '82
Magnalium, 86 Al, 13 Mg	24	St. '01	Gutta-percha	50.7	Vn. '02
Nickel steel, * 10% Ni	13.0	N.P.L.	Ice, -10° to 0°	25.1	B. '88
„ „ 20% „	19.5	N.P.L.	Iceland spar, axis	-5.6	B. '88
„ „ 30% „	12.0	N.P.L.	„ „ „ axis	—	—
„ „ 36% „	0.9	N.P.L.	Marble, white Carrara, 15°, 1.4 to 4°	3.5	N.P.L.
(Invar †)	6.0	N.P.L.	„ black	4.4	—
„ „ 40% „	9.7	N.P.L.	Masonry	7	—
„ „ 50% „	12.5	N.P.L.	Paraffin wax, 0°-40°	6.110	—
„ „ 80% „	16.8	B. '88	Porcelain, Berlin	2.8	S. '03
Phosphor bronze, 97.6 Cu, 2 Sn, 2 P	8.7	B. '88	„ „ 0°-100°	3.1	H.G. '01
Platinum-iridium, 90 Pt, 10 Ir ‡	15	—	„ Bayeux	3.4	Bd. '00
Platinum-silver, 33 Pt, 67 Ag	25	Sm.	„ „ 0°	2.5	T. '02
Solder, 2 Pb, 1 Sn, 50°	19.3	Sm.	Portland stone	6.3	—
Speculum metal, 68 Cu, 32 Sn	19	DI.	Quartz (crystal), axis	7.5	B. '88
Type metal, c. 135°	9.5	N.P.L.	„ „ „ axis	13.7	B. '88
Miscellaneous—			Silica (fused), -80° to 0°	2.2	S. '07
Brick (Egyptian)	14	—	„ „ 0° to 30°	4.2	C. '03
Cement and concrete, 10 to 77	77	—	„ „ 0° to 100°	5.0	S. '07
Ebonite	64.10	F. '68	„ „ 0° to 1000°	5.4	R. '10
Fluor spar, CaF ₂	8.5	Sc.	Sandstone	12	—
Glass, soft, 68 SiO ₂ , 14 Na ₂ O, 7 CaO	9.7	Sc.	Slate	10	—
„ hard, 64 SiO ₂ , 20 K ₂ O, 11 CaO			Woods (1) along grain—		
			Beech; mahogany	6.3	Vl. '68
			Oak; pine	6.5	Vl. '68
			(2) across grain—		
			Beech	60	Vl. '68
			Mahogany	40	Vl. '68
			Pine	34	Vl. '68

* See Guillaume's "Les Applications des Aciers au Nickel," 1904. † Invar is obtainable in three qualities, with a range of coefficients of $(-3 \text{ to } +2.5) \times 10^{-6}$ at ordinary temperatures. ‡ Used for international prototype metre (see p. 3). § Used for Imperial Standard Yard (see p. 4). B. Benoît; Bd. Bedford; C. Chappuis; D. Dittenberger; DI. Daniell; F. Fizeau; H. Hagen; H.D. Holborn and Day; H.G. Holborn and Grüneisen; M. Matthiessen; N.P.L. National Physical Laboratory; Pf. Pfaff; R. Randall; Ru. Russner; S. Scheel; Sc. Schott; Sm. Smeaton; St. Stadthagen; T. Tutton; T.S.S. Thiesen, Scheel, and Sell; V. Voigt; Vl. Villari; Vn. Vincent.

COEFFICIENTS OF CUBICAL EXPANSION OF GASES

The volume coefficient, α , at constant pressure is defined by $v_t = v_0(1 + \alpha t)$; the pressure coefficient, β , at constant volume is defined by $p_t = p_0(1 + \beta t)$, where v_t and p_t are the volume and pressure respectively corresponding to t° , the initial volume and pressure (v_0, p_0) being measured at 0°C . The values of both α and β depend on the initial pressure of the gas. If a gas obeys Boyle's law exactly, $\alpha = \beta$.

Comparison of rarefied gas, H_2 and absolute temperature scales.—

By graphically or otherwise extrapolating α and β to zero pressure, they become equal (as we should expect, for rarefied gases should behave as ideal gases and obey Boyle's law), and we may write $\alpha = \beta = \gamma$. For example, Berthelot finds from Chappuis' data—

For H_2 , mean $\gamma = .00366207 = 1/273.07$ (see p. 44)

N₂ $\gamma = .00366182 = 1/273.09$ (see p. 44)

Kelvin's absolute temperature scale agrees with the ideal gas scale, and therefore with the rarefied gas scale. Now, as will be seen below, β for $H_2 = \gamma$ very nearly, and thus the constant-volume hydrogen scale of temperature may justifiably be taken as closely approximating to the thermodynamic scale (see also p. 44).

(See Young's "Stoichiometry"; and Berthelot and Chappuis, *Trav. et Mém. du Bur. Intl.*, 1907.)

Gas.	Temp.	p_0 .	α	Obs.	Gas.	Temp.	p_0 .	β	Obs.
AT CONSTANT PRESSURE.					AT CONSTANT VOLUME.				
Air	C.	cm. Hg.			Air	C.	cm. Hg.		
	0°-100°	100° I	.0036728	C., 1903		—	.58	.0037666	M., 1892
	0-100	76	3671	R., 1847		—	1°32	37172	"
H₂	0-100	100	36600	C., 1903		—	10°0	36630	"
"	0-100	76	3661	R., 1847		—	17-24	36513	R., 1847
"	0-100	76	36609	R. M.		—	76	36650	"
N₂	0-100	100	367313	C., 1903		0°-100°	100° I	36744	C., 1903
"	0-100	139	367750	C., 1903		—	200	3690	R., 1847
"	—	200 atm.	434	A., 1890		—	2000	3887	"
"	—	1000 "	218	A., 1890		0-1067	23	36643	J. P.
O₂	—	100 "	486	A., 1890	H₂	0-100	52	36626	T. J., '02
CO	—	76	3669	R., 1847	"	0-100	70	366255	"
CO₂	0-20	51°8	37128	C., 1903	"	0-100	100	366256	C., 1903
"	0-40	"	37100	"	"	0-100	109	36627	O., 1908
"	0-100	"	37073	"	N₂	0-100	53	36683	C., 1903
"	0-20	99°8	37602	"	"	0-100	79	36718	"
"	0-40	"	37536	"	"	0-100	100	367440	"
"	0-100	"	37410	"	O₂	0-100	66	36738	M. N., '03
"	0-20	"	37972	"	"	0-1067	18-23	36652	J. P.
"	0-40	137°7	37906	"	He	0-100	52	36627	T. J., '02
"	0-100	"	37703	"	"	0-100	70	366255	"
"	0-100	"	37282	R. M.	"	0-100	100	36616	O., 1908
N₂O	—	76	3719	R., 1847	A	—	51°7	3668	K R., 96
NH₃	0-50	76/15°	3854	P. D., '06	CO	0-100	76	3667	R., 1847
SO₂	—	76	3903	R., 1847	"	0-1067	23	36648	J. P.
					CO₂	0-100	51°8	36981	C., 1903
					"	0-20	99°8	37335	"
					"	0-100	99°8	37262	"
					"	0-1067	24	36756	J. P.
					N₂O	—	76	3676	R., 1847
					SO₂	—	76	3845	R., 1847

COEFFICIENTS OF CUBICAL EXPANSION OF LIQUIDS

As with solids (see p. 52), if the temperature interval is not large, a linear equation $v_t = v_0(1 + \alpha t)$ may be employed to show the relation between the volume (v) of a liquid and its temperature (t). The mean coefficient (α) thus defined increases in general with the temperature. The values of α subjoined are per ° C., and for a range round 18° C. unless otherwise specified.

Liquid.	Temp. range.	Mean Coefficient from 0° C. to t ° C.	Observer.
Water (see p. 22 and below)	H scale. 17 to 40	$\cdot 0_{813019}/(t) - \cdot 0_{65769} + \cdot 0_{86797}t - \cdot 0_{7336}t^2$	Chappuis, '97
	17 to 100	Density = $1 - \frac{(t - 3982)^2}{466,700} \cdot t + 67 \cdot 365 - t$	Thiesen, '03
	Mercury 24 to 299	$\cdot 00018179 + \cdot 0_{9175}t + 0_{10351}t^2$	Regnault, '47 (Broch)
	0 to 100	$\cdot 00018169 - \cdot 0_{2817}t + 0_{9115}t^2$	Chappuis, '07
Mercury (see p. 22)	-10 to 300	$\cdot 000180555 + \cdot 0_{71244}t + \cdot 0_{10254}t^2$	{ Callendar & Moss, 1911 Donaldson, '12
	0 to 180	$\cdot 000181385 + \cdot 0_{9770}t + \cdot 0_{1018318}t^2$	

Liquid.	α	Liquid.	α	Liquid.	α	Liquid.	α
Acetic acid .	$\times 10^{-6}$ 107	Ether, ethyl .	$\times 10^{-6}$ 163	Pentane . .	$\times 10^{-6}$ 159	Water, 60-80	$\times 10^{-6}$ 58.7
Alcohol, me. .	122	Ethyl bromide	137	Toluene . .	109	Solutions—	
„ ethyl . .	110	Glycerine . .	53	Turpentine .	94		
„ amyl . .	93	Mercury (see above)		Xylol (m) .	101		
Aniline . .	85	Methyl iodide	121	Water, 5°-10°	5.3		
Benzene . .	124	Oil, olive . .	70	„ 10-20	15.0		
CS ₂	121	„ paraffin . .	90	„ 20-40	30.2	CaCl ₂ , 5.8% .	25.0
Chloroform .	126	„ „ 20°-199°	110	„ 40-60	45.8	„ 40.9% .	45.8
						NaCl, 26% .	43.6
						H ₂ SO ₄ , 100% .	57

MECHANICAL EQUIVALENT OF HEAT

Joule's equivalent, J , is here given as the number of ergs equivalent to a calorie, i.e. the heat required to raise 1 gram of water through 1° C. at some specified temperature. The **15° calorie** is about 1 part in 1000 greater than the **20° calorie**. (See p. 56.)

See Griffith's "Thermal Measurement of Energy," 1901.

Observer.	Calorie.	Ergs.	Observer.	Calorie.	Ergs.
	N. scale	$\times 10^7$		N. scale	$\times 10^7$
Joule, 1843	20° C.	4.169	Bousfield, <i>Phil. Trans.</i> , 1911	20° C	4.175
Rowland, 1878	20°	4.180	Crémieu & Rispail, 1908	0°	4.185
Griffiths, 1893	20°	4.184	Reynolds & Moorby, 1897	Mean	4.184
Schuster and Gannon, 1894	20°	4.181	Barnes, 1909 (deduced)	Mean	4.185
Callendar and Barnes, 1899	20°	4.180			

SPECIFIC HEATS

SPECIFIC HEAT OF WATER

Callendar and Barnes (*Phil. Trans.*, 1902) used an electrical method of determining the temperature variation of the specific heat of water. The specific heats below are reduced by Callendar ("Ency. Brit." Art. "Calorimetry") from their results; they are relative to the specific heat at 20° C. on the C.P. nitrogen scale. The 20° calorie (see pp. 5 and 55) is adopted as 4'180 joules = 4'180 × 10⁷ ergs, being the mean of the results of Rowland (1879) and of Reynolds and Moorby (reduced), each of whom used a mechanical method of determining "J." Thus the values of J below do not rest on the values attributed to the electrical standards employed. The specific heat of water is a minimum at 37'5° C., according to Callendar and Barnes.

The 15° calorie (according to Barnes, *Proc. Roy. Soc.*, 1909) = 4'184 joules, assuming the e.m.f. of the Clark cell at 15° C. = 1'4330 international volts.

The mean calorie (= $\frac{1}{100}$ of heat required to raise 1 gram of water from 0° to 100° C.) = 4'185 joules (Barnes, 1909); = 4'184 joules (Reynolds and Moorby, 1897, corrected by Smith).

Temp.	Specific heat.	Joules.	Temp.	Specific heat.	Joules.	Temp.	Specific heat.	Joules.
-5° C.	1'0158	4'246	45° C.	'9983	4'173	95° C.	1'0063	4'206
0	1'0094	4'219	50	'9987	4'175	100	1'0074	4'211
5	1'0054	4'202	55	'9992	4'177	120	1'0121	4'231
10	1'0027	4'191	60	1'0000	4'180	140	1'0176	4'254
15	1'0011	4'184	65	1'0008	4'183	160	1'0238	4'280
20	1'0000	4'180	70	1'0016	4'187	180	1'0308	4'309
25	'9992	4'177	75	1'0024	4'190	200	1'0384	4'341
30	'9987	4'175	80	1'0033	4'194	220	1'0467	4'376
35	'9983	4'173	85	1'0043	4'198			
40	'9982	4'173	90	1'0053	4'202			

SPECIFIC HEAT OF MERCURY

In terms of the gram calorie at 15°·5 on the const. vol. H. scale. (Barnes and Cooke, *Phys. Rev.*, 15, 1902.) Mercury has a minimum specific heat at 146° C. (Barnes, *Brit. Ass. Rep.*, 1909.)

Temp.	0° C.	20°	40°	60°	80°	100°	200°
Specific heat	'0335	'0333	'0331	'0329	'0328	('0327)	('032)

SPECIFIC HEATS OF THE ELEMENTS

For gases, see p. 58. (See Waterman, *Phys. Rev.*, 1896.)

Substance.	Temperature.	Sp. heat.	Observer.	Substance.	Temperature.	Sp. heat.	Observer.
Aluminium .	-182° to 15°	'168	Tilden, 1903	Bromine, liqd.	13° to 45°	'107	Andrews, '48
	15 to 185	'219	"	Cadmium *	-186 to -79	'050	Behn, 1900
	600	'282	Richards, '93	" pure	18 to 99	'055	Voigt, 1893
Antimony . .	-186 to -79	'0462	Behn, 1900	Cæsium . .	0 to 26	'048	E. & G., 1900
	17 to 92	'0508	Gaede, 1902	Calcium . .	-185 to 20	'157	N. & B., 1906
Arsenic, cryst.	21 to 68	'083	B. & W., 1868		0 to 100	'149	Be., 1906
" amorph.	21 to 65	'076	"	Carbon—			
Barium . . .	-185 to 20	'068	N. & B., 1906	Gas carbon .	24 to 68	'204	B. & W., 1868
Beryllium . .	0 to 100	'425	N. & P., 1880	Charcoal . .	0 to 24	'165	H.F. Weber, '75
Bismuth . . .	-186	'0284	Giebe, 1903	"	0 to 224	'238	"
	22 to 100	'0304	W., 1896	Graphite . .	-50	'114	"
Boron, amor.	0 to 100	'307	M. & G., 1893	"	11	'160	"
Bromine, solid	-78 to -20	'084	Regnault, '49	"	202	'297	"

* Contained Fe and Zn.

SPECIFIC HEATS OF THE ELEMENTS (*contd.*)

Substance.	Temperature.	Sp. heat.	Observer.	Substance.	Temperature.	Sp. heat.	Observer.
Carbon (<i>contd.</i>)				Palladium . .	-186° to 18°	·053	Behn, 1898
Graphite . .	977° C.	·467	H.F. Weber, '75		18 to 100	·059	"
Diamond . .	-50	·064	"	Phosphorus—			
" . .	11	·113	"	" yellow	-78 to 10	·17	Regnault, 1849
" . .	206	·273	"		13 to 36	·202	Kopp, 1864
" . .	985	·459	"	" liquid	49 to 98	·205	Person, 1847
Cerium . .	0 to 100	·045	H., 1876	" red . .	15 to 98	·17	Regnault, 1853
Chlorine, liq.	0 to 24	·226	Knietsch	Platinum . .	-186 to 18	·0293	Behn, 1898
Chromium . .	-200	·067	Adler, 1903		18 to 100	·0324	"
(1·4% Fe & Si)	0	·104	"		1230	·0461	Tilden, 1903
	100	·112	"	Potassium . .	-78 to 23	·166	Schütz, 1892
	400	·133	"	Rhodium . .	10 to 97	·058	Regnault, 1862
Cobalt . . .	-182 to 15	·082	Tilden, 1903	Ruthenium . .	0 to 100	·061	Bunsen, 1870
	15 to 100	·103	"	Selenium, cryst.	22 to 62	·084	B. & W., 1868
	15 to 630	·123	"	" amorph.	18 to 38	·095	"
Copper . . .	-192 to 20	·0798	Schmitz, 1903	Silicon, cryst.	-185 to 20	·123	N. & B., 1906
	20 to 100	·0936	"		57	·183	H.F. Weber, '75
	900	·118	Le Verrier, '92		232	·203	"
Didymium . .	0 to 100	·046	H., 1876	Silver . . .	-186 to -79	·0496	Behn, 1900
Gallium, solid	12 to 23	·079	B., 1878		15 to 100	·056	B. & S., 1895
" liquid	12 to 119	·080	"		427	·059	Tilden, 1903
Germanium . .	0 to 100	·074	N. & P., 1887	Sodium . . .	-185 to 20	·234	N. & B., 1906
Gold . . .	-185 to 20	·035	N. & B., 1906		10	·297	Bernini, 1906
	18 to 99	·0303	Voigt, 1893		128	·333	"
Indium . . .	0 to 100	·057	Bunsen, 1870	Sulphur—			
Iodine . . .	9 to 98	·054	Regnault, 1840	" rhombic	17 to 45	·163	Kopp, 1865
Iridium . . .	-186 to 18	·0282	Behn, 1898	" liquid . .	119 to 147	·235	Person, 1847
	18 to 100	·0323	"	Tantalum . .	-185 to 20	·033	N. & B., 1906
Iron . . .	-192 to 20	·089	Schmitz, 1903		58	·036	v. Bolton, 1905
	20 to 100	·119	"	Tellurium, crys.	15 to 100	·048	Fabre, 1887
	225	·137	Stücker, 1905	Thallium . .	-192 to 20	·0300	Schmitz, 1903
	0 to 1100	·153	Harker, 1905		20 to 100	·0326	"
Lanthanum . .	0 to 100	·045	H., 1876	Thorium . .	0 to 100	·028	Nilson, 1883
Lead . . .	-192 to 20	·0293	Schmitz, 1903	Tin . . .	-186 to -79	·0486	Behn, 1900
	20 to 100	·0305	"		19 to 99	·0552	Voigt, 1893
	300	·0338	Naccari, 1888	" molten . .	240	·064	Spring, 1886
Lithium . . .	0 to 19	·837	Be., 1906	Titanium . .	-185 to 20	·082	N. & B., 1906
	0 to 100	·1093	"		0 to 100	·113	N. & P., 1887
Magnesium . .	-186 to -79	·189	Behn, 1900		0 to 440	·162	"
	18 to 99	·246	Voigt, 1893	Tungsten . .	-185 to 20	·036	N. & B., 1906
	225	·281	Stücker, 1905		20 to 100	·034	Gin, 1908
Manganese . .	14 to 97	·122	Regnault, 1862	Uranium . .	11 to 98	·062	Regnault, 1840
Mercury . . .	See preceding page.				0 to 98	·028	Blümcke, 1885
Molybdenum .	-185 to 20	·063	N. & B., 1906	Vanadium . .	0 to 100	·115	Mache, 1897
	15 to 91	·072	D. & G., 1901	Zinc . . .	-192 to 20	·084	Schmitz, 1903
Nickel . . .	-186 to 18	·086	Behn, 1898		20 to 100	·093	"
	18 to 100	·109	"		300	·104	Naccari, 1888
Osmium . . .	19 to 98	·031	Regnault, 1862	Zirconium . .	0 to 100	·066	M. & D., 1873

B., Berthelot; Be., Bernini; B. & S., Bartoli & Stracciati; B. & W., Bettendorff & Wüller; D. & G., Defacqz & Guichard; E. & G., Eckardt & Graefe; H., Hillebrand; M. & D., Mixer & Dana; M. & G., Moissan & Gautier; N. & B., Nordmeyer & Bernoulli; N. & P., Nilson & Pettersson; W., Waterman.

SPECIFIC HEATS

SPECIFIC HEATS OF GASES AND VAPOURS

The values at const. pressure are, unless otherwise stated, all at atmospheric pressure. The specific heats given are calories per gram of gas per degree C. at the temp. stated.

Gas.	Temp.	Sp. ht.	Observer.	Gas.	Temp.	Sp. ht.	Observer.
AT CONSTANT PRESSURE (c_p)				Ammonia, NH_3 .	23-100°	.520	Wiedemann
Air (dry)	20° C.	.2417	Swann, 1909	Nitrous oxide, N_2O	26-103	.213	1876
" "	100	.2430	" "	Nitric oxide, NO	13-172	.232	Regnault, '62
" "	20-440	.2366	H. & A., 1905	N. peroxide, NO_2	27-67	.1625	B. & O., 188
" "	20-98	.2372	Witkowski,	H_2S	20-206	.245	Regnault, '62
" "	-102-17	.2372	" 1896	CS_2	86-190	.160	" "
" " 70 atmos.	-50	.312	" "	Methane, CH_4 . .	—	.591	Lussana, '94
Argon	20-90	.123	D., 1897	Ethylene C_2H_4 . .	—	.404	" "
Hydrogen	—	.3402	Lussana, 1894	Benzene, C_6H_6 . .	34-115	.299	Wiedemann
" " 30 atmos.	—	.3788	" "	Chloroform CHCl_3 .	27-118	.144	1877
Nitrogen	0	.2350	* H. & H., '07	Me. alcohol CH_3O .	101-223	.458	Regnault, '62
" (liq.)	-200	.43	Alt, 1904	Et. alcohol $\text{C}_2\text{H}_5\text{O}$.	108-220	.453	Regnault, '62
Oxygen	20-440	.2419	H. & A., 1905	" ether (C_2H_5) $_2\text{O}$.	25-111	.428	W., 1876
" "	20-800	.2497	" "	Turpentine, $\text{C}_{10}\text{H}_{16}$	179-249	.506	Regnault, '62
" (liq.)	-190	.347	Alt, 1904				
Chlorine	16-343	.115	Strecker, '81	AT CONSTANT VOLUME (c_v)			
Bromine	19-388	.055	" '82	Air, † 1 atmos . . .	0°	.1715	Joly, 1891
Iodine	206-377	.034	" "	Hydrogen ‡ . . .	c. 50	.2402	" "
Carbon monoxide .	23-99	.242	W., 1876	Carbon dioxide § .	c. 55	.1650	" 1894
" dioxide	0	.2010	* H. & H., '07	Argon	0-2000	.0746	Pier, 1909
" "	20	.2020	Swann, 1909	Nitrogen 	0	.175	" "
" "	100	.221	" "	Water vapour . . .	100	.340	" "
" " 30 atmos.	2670	.2670	Lussana, '94				
Water vapour . .	100	.4652	* H. & H., '07				

B. & O., Berthelot & Ogier; D., Dittenberger; H. & A., Holborn & Austin (Reichsanstalt); W., Wiedemann.
 * H. & H., Holborn & Henning { Nitrogen (0-1400°), $c_p = .2350 + .000019t$ } Mean specific
 (Reichsanstalt). { CO_2 (0-1400°), $c_p = .2010 + .0000742t - .0.18t^2$ } heats between
 { Water vapour (100-1400°), $c_p = .4669 - .0000168t + .0.44t^2$ } 0° and t° C.

† Air, $c_v = .1715 + .02788\rho$ where ρ is the density (gm./c.c.). § CO_2 , $c_v = .165 + .2125\rho + .34\rho^2$, ρ being density.
 ‡ H, c_v diminishes with increasing density and falling temp. || N, $c_v = .175 + .00016t$, t being the temp.

RATIO OF THE SPECIFIC HEATS FOR GASES AND VAPOURS

γ = the ratio of the specific heat at constant pressure to that at constant volume. γ is usually determined directly by some method involving an adiabatic expansion, such as the determination of the velocity of sound in the gas. From a knowledge of either (1) the pressure or (2) the temperature immediately following an adiabatic expansion (Clément and Desormes, Lummer and Pringsheim's methods respectively), γ can be deduced from $pV^\gamma = \text{const.}$, or $\theta V^{\gamma-1} = \text{const.}$ (See Capstick, "Science Progress," 1895; and Moody, *Phys. Rev.*, Ap., 1912.)

Gas.	Temp.	γ	Observer.	Gas.	Temp.	γ	Observer.
Monatomic gases				Air (dry)	0°	1.402	Koch, 1907
Helium	0° C.	1.63	B. & G., 1907	" "	0	1.402	F., 1908
Argon	0	1.667	Niemeyer, '02	" "	500	1.399	" "
Neon	19	1.642	Ramsay, 1912	" "	900	1.39	Kalähne, '03
Krypton	19	1.689	" "	" "	-79.3	1.405	Koch, 1907
Xenon	19	1.666	" "	" " 200 }	0	1.828	" "
Mercury vapour .	310	1.666	K. & W., 1876	" " atmos. }	-79.3	2.333	" "
Diatomic gases—				Hydrogen	—	1.419	Hartmann, '0
Air (dry)	5-14	1.402	L. & P., 1898	" "	4-16	1.408	L. & P., 1898
" "	0	1.401	Stevens, 1905	Nitrogen	—	1.41	Cazin, 1862
" "	15	1.401	Makower, '03	Oxygen	5-14	1.400	L. & P., 1898
" "	—	1.414	Hartmann, '02	Carbon monoxide .	—	1.401	Leduc, 1898
				Nitric oxide, NO .	—	1.394	Masson

B. & G., Behn & Geiger; F. Fürstenau; K. & W. Kundt & Warburg; L. & P., Lummer & Pringsheim.

RATIO OF THE SPECIFIC HEATS FOR GASES AND VAPOURS (*contd.*)

Gas.	Temp.	γ	Observer.	Gas.	Temp.	γ	Observer.
Triatomic gases				Acetylene, C_2H_2	—	1'26	M. & F., 1897
Ozone	—	1'29*	Jacobs, 1905	Ethylene, C_2H_4	—	1'264	Capstick, '95
Water vapour . . .	100° (?)	1'305	Makower, '03	Benzene, C_6H_6	20°	1'40	Pagliani, '96
Carbon dioxide . .	4-11	1'300	L. & P., 1898	"	99'7	1'105	Stevens, '02
" "	—	1'306	Hartmann, '05	Chloroform, $CHCl_3$	24-42	1'110	Müller, 1883
" "	500	1'26	F., 1908	"	99'8	1'150	Stephens, '02
Ammonia, NH_3 . .	—	1'336	Leduc, 1898	CCl_4	—	1'130	Capstick, '95
Nitrous oxide, N_2O	—	1'324	"	Me. alcohol . . .	99'7	1'256	Stevens, '02
Nitrogen $\{ N_2O_4$. .	20°	1'172	Natanson, '85	" bromide . . .	—	1'274	Capstick, '93
peroxide $\{ NO_2$. .	150	1'31	"	" chloride . . .	19-30	1'279	" "
H_2S	—	1'340	Capstick, '95	" iodide . . .	—	1'286	" "
CS_2	—	1'239	"	Et. alcohol . . .	53	1'133	Jaeger, 1889
Sulphur dioxide. {	16-34	1'26	Müller, 1883	" "	99'8	1'134	Stevens, '02
	500	1'2	F., 1908	" bromide . . .	—	1'188	Capstick, '93
Polyatomic gases				" chloride . . .	22'7	1'187	" "
Methane, CH_4 . .	—	1'313	Capstick, '93	" ether	12-20	1'024	Low, 1894
Ethane, C_2H_6 . .	—	1'22	{ Daniel &	" "	99'7	1'112	Stevens, '02
Propane, C_3H_8 . .	—	1'130	{ Pierron, '99	Acetic acid . . .	136'5	1'147	" "

* Extrapolated; F., Fürstenau; L. & P., Lummer & Pringsheim; M. & F., Maneuvrier and Fournier.

SPECIFIC HEATS OF VARIOUS BODIES

In most cases, the specific heats given must only be regarded as rough average values.

Substance.	Temp.	Sp. ht.	Substance.	Temp.	Sp. ht.	Substance.	Temp.	Sp. ht.
Alloys—			Ether, ethyl . .	18°	'56	Glass, Jena 16'''†	18°	'19
Brass, red . . .	0	'090	Glycerine . . .	18-50	'58	" Jena 59'''†	18	'19
" yellow . . .	0	'088	Oil, olive . . .	7	'47	Granite	20-100	{ '19 to '20
Eureka	18	'098	" paraffin . . .	20-60	{ '51 to '54	Ice	-21 to -1	{ '502
(Constantan)			Sea-water . . .	17	'94	Indiarubber . .	15-100	{ '27 to '48
German silver .	0-100	'095	Toluene	18	'40	Marble, white .	18	{ '21 to '22
Liquids—			Turpentine . . .	18	'42	Paraffin wax . .	0-20	'69
Alcohol, amyl .	18	'55	Miscellaneous—			Porcelain ‡ . .	15-1000	'255
" ethyl . . .	0	'547	Asbestos . . .	20-100	'20	Quartz, SiO_2 .	0	'174
" "	40	'648	Basalt	20-100	{ '20 to '24	" " " " . . .	350	'279
" methyl . . .	12	'601	Ebonite	20-100	'33	Rock salt, $NaCl$	18	'21
Aniline * . . .	15	'514	Fluorspar, CaF_2	30	'21	Sand	20-100	'19
Benzene	10	'340	Glass, crown . .	10-50	'16	Silica (fused) §.	15-200	'200
" "	40	'423	" flint	10-50	'12	" "	15-800	'248
Brine, density = 1'2	-20	'69						
(Harker) . . .	0	'71						
	15	'72						

* Griffiths, *Phil. Mag.*, 1893.

† See p. 74.

‡ Harker, 1905.

§ Greenwood, 1911.

LATENT HEATS

LATENT HEAT OF FUSION

The number of gram calories required to convert 1 gram of substance from solid into liquid without change of temperature.

ICE

Temp.	Lt. ht.	Observer, etc.
-6.5° C.	cal. 76.03	Pettersson, 1881.
0	79.59	Regnault, 1843, corrected.
0	80.02	Bunsen, 1870, with ice calorimeter.
0	79.77	Smith, <i>Phys. Rev.</i> , 1903 (in terms of 15° calorie = 4.184 joules, taking Clark cell = 1.433 volts at 15° C.).

VARIOUS SUBSTANCES

Substance.	Temp.	Lt. ht.	Substance.	Temp.	Lt. ht.	Substance.	Temp.	Lt. ht.
Elements—	°C.	cal.		°C.	cal.		°C.	cal.
Aluminium . . .	657	77	Platinum . . .	1750	27	NaNO ₃ . . .	311	63
Bismuth . . .	269	13	Potassium . . .	62	16	KNO ₃ . . .	339	47
Cadmium . . .	321	14	Silver . . .	960	22	H ₂ SO ₄ . . .	10.3	24
Copper . . .	—	43	Sulphur . . .	115	9	Acetic acid . . .	4	44
Lead . . .	327	5	Tin . . .	232	14	Benzene . . .	5.4	30
Mercury . . .	—	3	Zinc . . .	418	28	Glycerine . . .	13	42
Palladium . . .	1550	36	Compounds—			Naphthalene . . .	80	35
Phosphorus . . .	44	5	NH ₃ . . .	-75	108	Xylene . . .	16	39

LATENT HEAT OF VAPORISATION

Latent heats are given as the number of gram calories required to convert 1 gram of substance from liquid into vapour without change of temperature. The latent heat of vaporisation vanishes at the critical temperature.

Trouton's Rule.—The latent heat of vaporisation of 1 gramme molecule of a liquid divided by the corresponding boiling point (on the absolute scale) is a constant (C). $C = 21$ for substances of which both liquid and vapour are unassociated. If the liquid is associated, $C > 21$ (e.g. water, $C = 26$); if the vapour is associated, $C < 21$ (e.g. acetic acid, $C = 15$). [See Nernst's "Theoretical Chemistry."]

STEAM

Regnault's equation connecting latent heat and temperature takes no account of the temperature variation of the specific heat of water (see p. 56). The equation gives values which are too large at low temperatures. The equations of Griffiths, Henning, and Smith have been reduced and are here expressed in terms of the **15° calorie** = 4.184 joules. Griffiths' and Smith's results rest further on an attributed value of 1.433 volts for the e.m.f. of the Clark cell at 15° C.

See also next page.

[The critical temp. of water is about 365° C.]

Observer.	Temp. range of expts.	Latent heat L_t at $t^\circ \text{C}$.
Regnault, 1847 . .	63°–194° C.	$L_t = 606.5 - .695t$
Griffiths, 1895 . .	30° and 40°	$L_t = 598.0 - .605t$
Henning, <i>Ann. d. Phys.</i> , 1906, . .	30°–100°	$L_t = 599.4 - .60t$, to .3 %
1909 . . .	100°–180°	or $L_t = 94.3 (365 - t)^{.312}$, to .1 %
Smith, <i>Phys. Rev.</i> , 1907 . .	14°–40°	$L_t = 538.97 - .6428(t - 100) - .03834(t - 100)^2$
		$L_t = 597.2 - .580t$

LATENT HEAT OF STEAM (contd.)

In terms of 15° calorie	Regnault, 1847.	Griffiths, 1895.	Joly, 1895.	Callendar, *	Dieterici, 1905.	Henning, 1906.	Smith, 1911.	Richards & Matthews, 1911.
L_0 . .	606†	598†	—	595†	596·0†	599†	—	—
L_{100} . .	537	537·5†	540§	540	538·9	539·4	540·5	538·0

* From sp. ht. of steam experiments and total heat formula.

† Extrapolated.

‡ Reduced to mean calories (4·185 joules); Clark cell = 1·433 volts.

§ By comparing L_{100} (by steam calorimeter) with the mean specific heat of water between 12° and 100°.

Callendar and Barnes' specific heat has been used (p. 56).

|| Carlton-Sutton, 1917.

LATENT HEATS OF VAPORISATION OF VARIOUS SUBSTANCES

The values below are for pure substances, and are due to Young, *Proc. Roy. Dublin Soc.*, 1910. The precise calorie employed is not stated.

Temp.	SnCl_4	CCl_4	Pent- ane (n).	Methyl	Ethyl	Propyl	Ethyl ether.	Methyl	Ethyl	Propyl	Acetic acid.	Ben- zene.
				Alcohol.				Acetate.				
	cal.	cal.	cal.	cal.	cal.	cal.	cal.	cal.	cal.	cal.	cal.	cal.
C.	—	—	—	289·2	220·9	—	92·52	—	—	—	—	—
0°	—	—	—	284·5	220·6	—	87·54	—	—	—	84·05	—
20	—	—	—	277·8	218·7	—	82·83	—	—	—	87·02	—
40	—	—	84·31	269·4	213·4	—	78·44	98·59	—	—	89·69	—
60	—	—	80·07	259·0	206·4	173·0	73·50	94·07	—	79·80	91·59	95·45
80	—	46·00	75·33	246·0	197·1	164·0	68·42	88·39	85·78	76·33	92·32	91·41
100	31·76	44·15	69·94	232·0	184·2	153·0	62·24	82·87	82·15	71·84	94·38	86·58
120	30·54	42·08	64·48	216·1	171·1	142·4	55·93	76·83	72·24	67·66	91·83	82·82
140	29·12	39·92	56·58	198·3	156·9	129·0	46·07	69·96	65·91	62·80	89·63	78·94
160	27·69	37·95	47·42	177·2	139·2	116·3	31·87	61·00	59·87	57·23	87·71	74·62
180	26·29	35·40	35·01	151·8	116·6	102·2	19·38†	50·56	52·71	50·78	85·55	68·81
200	24·57	32·61	24·68*	112·5	88·2	85·3	—	34·87	42·63	42·40	82·02	62·24
220	22·82	29·45	—	84·5†	40·3	63·4	—	20·99§	27·17	30·70	78·18	54·11
240	20·86	25·56	—	—	—	33·5	—	—	12·03	11·73	72·26	43·82
260	18·50	20·07	—	—	—	—	—	—	—	—	63·48	27·43
280	15·60	10·43	—	—	—	—	—	—	—	—	—	—
Crit. temp.)	318°·7	283°·1	197°·2	240°	243°·1	263°·7	193°·8	233°·7	250°·1	276°·2	321°·6	288°·5

* At 190°.

† At 230°.

‡ At 190°.

§ At 230°.

|| At 249°.

¶ At 275° C.

Substance.	Temp	Lt. ht.	Substance.	Temp.	Lt. ht.	Substance.	Temp.	Lt. ht.
	C.	cal.		C.	cal.		C.	cal.
Mercury . .	358°	68	Liquid N_2O . .	-20°	67	Chloroform . .	61°	58
Sulphur . .	316	362	" NH_3 . .	—	341	Et. bromide . .	38	60
Phosphorus . .	287	130	" CO_2 . .	0	57	" propionate . .	100	79
Liquid H_2 . .	—	123	" SO_2 . .	22	32	" iodide . .	71	47
" O_2 . .	-188	58	" CS_2 . .	-10	96	" formate . .	50	98
" N_2 . .	—	50	Me. formate . .	32·5	110·5	Am. alcohol . .	131	120
" air . .	—	c. 50	" iodide . .	42	46	Aniline . .	—	104
" Cl . .	-22	67	Chloroform . .	0	67	Toluene . .	111	84
Bromine . .	58	46				Turpentine . .	159	70
Iodine . .	174	24						

THERMOCHEMISTRY

THERMOCHEMISTRY

In thermochemistry the conservation of energy is assumed in accordance with experiment, and consequently (1) if a cycle of chemical change takes place so that the final state of the reacting substances is identical with the initial, then as much heat is absorbed as is given out, *i.e.* the total heat of the reaction is zero; (2) the heat of reaction only depends on the initial and final states of the reacting substances, and not on the intermediate stages. The results below are affected by, but have not been corrected for, any changes in the accepted values of the atomic weights since the experiments were carried out.

MOLECULAR HEAT OF FORMATION

The **molecular heat of formation** (H.F.) is the heat liberated when the molecular weight in grams of a compound is formed from its elements. When the state of aggregation of an element or compound is not given, it is the state in which it occurs at room temperature and pressure. A minus sign before an H.F. means that heat is absorbed in the building up of the compound.

Unit—the gram calorie (at 15° to 20° C.) per gm. molecule of compound. Aq = solution in a large amount of water. The reactions are at constant pressure.

Example.—H.F. of $\text{CuSO}_4 = 183,000$; of $\text{CuSO}_4 \cdot \text{Aq} = 198,800$. \therefore the heat of solution of $\text{CuSO}_4 = 198,800 - 183,000 = 15,800$ cal. per gram mol.

(T., Thomsen, "Thermochemistry," trans. by Miss K. A. Burke; B., Berthelot, *Ann. d. Chim. et d. Phys.*, 1878; T.B., mean of both these observers' values. For organic compounds, see p. 64.

INORGANIC COMPOUNDS

Compound.	Mol. H.F. in calories.	Compound.	Mol. H.F. in calories.	Compound.	Mol. H.F. in calories.
Non-Metals	$\times 10^3$		$\times 10^3$		$\times 10^3$
HCl gas . .	22.0, T.	CO_2 from	97.3, B.T.	$\text{NH}_4\text{Cl} \cdot \text{Aq}$.	72.4
HCl. Aq . .	39.3, T.	amorph. C }		$(\text{NH}_4)_2\text{SO}_4$.	283, T.B.
HBr gas . .	8.4, T.	CO_2 from	94.3, B.	$(\text{NH}_4)_2\text{SO}_4 \cdot \text{Aq}$	280.6
HBr. Aq . .	28.6, T.	diamond }		$\text{NH}_4\text{OH} \cdot \text{Aq}$.	90, B.
HI gas . .	-6.1, T.B.	B_2O_3 ; amp. B.	273, B.	BaO	126, T.
HI. Aq . .	+13.2, T.B.	SiO_2 Aq; crys.	180, B.	$\text{Ba}(\text{OH})_2$. .	217, T.
HF	+38.5	As_2O_3 . . [Si]	155, T.	BaCl_2	197, T.
H_2O liq. .	68.4, T.	As_2O_5 . .	219, T.	$\text{BaCl}_2 \cdot \text{Aq}$. .	199.1, T.
" " . . .	69.0, B.	CCl_4 from	76, B.	Bi_2O_3	20
" gas . .	58.1, B.	diamond }		BiCl_3	91, T.
$\text{H}_2\text{O}_2 \cdot \text{Aq}$.	47.0	SbCl_3 solid .	91.4, T.	$\text{Cd}(\text{OH})_2$. . }	66, T.
H_2S from	2.7, T.	SbCl_3 liq. .	105, T.	$\text{Cd} + \text{O} + \text{H}_2\text{O}$ }	
rhombic S. }		CS_2 from		CdCl_2	93, T.
NH_3	12	diamond &	-19, B.	CdSO_4	222, T.
AsH_3 . . .	-36.7	rhombic S. }		$\text{CdSO}_4 \cdot 8/3 \text{H}_2\text{O}$	+2.66, T.
SbH_3 . . .	-87, B.	C_2N_2 gas }	-74, B.	on sol. in Aq }	
SiH_4 . . .	25	from diam. .		$\text{CdSO}_4 \cdot \text{Aq}$.	232.7, T.
SO_2 from	70	H_2SO_4 liq. .	193, T.	Cs_2O	100
rhombic S. }		$\text{H}_2\text{SO}_4 \cdot \text{Aq}$ }		CaO	131, T.
SO_3 liq. from	103	from rhombic	210, T.	" Moissan .	145
rhombic S. }		S.		$\text{Ca}(\text{OH})_2$. .	229
N_2O	-19	HNO_3 liq. .	41.6, B.	CaC_2	-7.25
NO	-21.6, T.	$\text{HNO}_3 \cdot \text{Aq}$.	49	CaCl_2	170, T.
N_2O_3 . . .	-21.4, B.	HCN gas }	-30.5	$\text{CaCl}_2 \cdot \text{Aq}$.	187.4, T.
$\text{NO}_2/22^\circ$. .	-1.7, B.	from diam. }		CaSO_4	318, T.
" /150° . .	-7.6, B.	HCN liq. .	-24.8	CaCO_3 . . .	270, T.
N_2O_5 liq. .	3.6, T.	H_3PO_4 liq. .	302	$\text{Ca}(\text{NO}_3)_2$. .	202, B.
P_2O_5 solid	369			Co	64
$\text{P}_2\text{O}_5 \cdot \text{Aq}$.	405	Metals—		CoCl_2	76.5, T.
CO from	29, T.	Al_2O_3	380, B.	$\text{CoSO}_4 \cdot 7\text{H}_2\text{O}$	234, T.
amorph. C. }		AlCl_3	161	$\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$	119, T.
CO from	26.1, B.	$\text{Al}_2(\text{SO}_4)_3 \cdot \text{Aq}$	880	CuO	37.2, T.
diamond }		NH_4Cl . . .	76.3, T.B.	CuCl_2	51.6

INORGANIC COMPOUNDS (contd.)

Compound.	Mol. H.F. in calories.	Compound.	Mol. H.F. in calories.	Compound.	Mol. H.F. in calories.
Metals (contd.)	$\times 10^3$		$\times 10^3$		$\times 10^3$
CuSO ₄ . . .	183, T.	MgCl ₂ . . .	151, T.	AgCl . . .	29.2, T.B.
CuSO ₄ .Aq . . .	198.8, T.	MgSO ₄ . . .	302, T.	Na ₂ O . . .	91 to 100
CuSO ₄ .5H ₂ O } on sol. in Aq. }	-275	MgSO ₄ .Aq . . .	322	NaHO . . .	102.3, T.B.
AuBr ₃ . . .	8.8, T.	MnO . . .	91	NaHO.Aq . . .	112.2, T.B.
AuCl ₃ . . .	23, T.	MnCl ₂ . . .	112	NaCl . . .	97.8, T.B.
FeO . . .	64.6	Hg ₂ O . . .	24.9, T.	NaNO ₃ . . .	111, T.B.
Fe ₂ O ₃ /400° . . .	196	HgO . . .	21.1	Na ₂ SO ₄ . . .	328.3, T.B.
Le Chatelier . . .		HgSO ₄ . . .	175	Na ₂ CO ₃ . . .	272, T.B.
FeSO ₄ .7H ₂ O . . .	240	HgCl . . .	31.3	SrO . . .	130, T.B.
FeSO ₄ .Aq . . .	236	HgCl ₂ . . .	53.2	Sr(OH) ₂ . . .	21.7, B.
FeCl ₃ . . .	96, T.	NiO . . .	59.7	SrCl ₂ . . .	185, T.B.
PbO . . .	50.3, T.	NiCl ₂ . . .	74.5, T.	SrCl ₂ .Aq . . .	196, T.
PbO ₂ . . .	62.4	NiSO ₄ .Aq . . .	22.9, T.	Tl ₂ O . . .	42.2, T.
PbCl ₂ . . .	83, T.	PtCl ₄ . . .	59.4	TlCl . . .	48.6, T.
PbSO ₄ . . .	216, T.	K ₂ O . . .	97	Tl ₂ SO ₄ . . .	221, T.
Pb(NO ₃) ₂ . . .	105.5	KHO . . .	104, B.T.	SnO . . .	70
Pb(NO ₃) ₂ .Aq . . .	97.9	KHO.Aq . . .	117, B.T.	SnCl ₂ . . .	81, T.
Li ₂ O . . .	140	KCl . . .	106, B.T.	SnCl ₄ . . .	128
LiOH . . .	111	KCl.Aq . . .	101.6, T.	ZnO . . .	85.4, T.
LiCl . . .	94, T.	KNO ₃ . . .	119, B.T.	ZnCl ₂ . . .	97.3, T.B.
LiCl.Aq . . .	102.4	K ₂ SO ₄ . . .	344, T.B.	Zn(NO ₃) ₂ .Aq . . .	132
Li ₂ SO ₄ . . .	334, T.	Ag ₂ O . . .	5.9, T.	ZnSO ₄ . . .	230.3, T.B.
LiNO ₃ . . .	112, T.	" . . .	7, B.	ZnSO ₄ .Aq . . .	248.7
MgO . . .	143, B.	AgNO ₃ . . .	28.7, T.B.	ZnSO ₄ .7H ₂ O } on sol. in Aq. }	-4.26
		AgNO ₃ .Aq . . .	23.3, T.		

MOLECULAR HEAT OF NEUTRALISATION

Unit—the gram calorie (at 15° to 20°) per gram molecule of base. Thus KOH.Aq + HCl.Aq = KCl.Aq + H₂O + 13,750 calories. Thomsen (= T.) observed at 18° to 20° C., and the final dilution was 3600 gms. (7200 for Na salts) per gm. mol. of base. Berthelot (= B.) used at least 2000 gms. of H₂O per 17 gms. of hydroxylyon, -HO.

Base.	HCl	HF	HNO ₃	HCN	$\frac{1}{2}$ H ₂ SO ₄	$\frac{1}{2}$ H ₂ CO ₃	1H ₃ PO ₄	1Oxalic.
	$\times 10^3$	$\times 10^3$	$\times 10^3$	$\times 10^3$	$\times 10^3$	$\times 10^3$	$\times 10^3$	$\times 10^3$
1NaOH . . .	13.74, T.; 13.7, B.	16.3, T.	13.7, T.; 13.5, B.	2.8	15.64, T.	10.1, T.; 10.2, B.	14.8, T.	13.8, T.
2NaOH . . .	—	—	—	—	31.38†, T.	20.2‡, T.	27.1*, T.	28.3, T.
1LiOH . . .	13.85, T.	16.4 †	—	2.93	15.64, T.	—	—	—
1KOH . . .	13.7, T.; 13.6, B.	16.1	13.8, T.	2.8, T.	15.7, T.B.	10.1, B.	—	13.8, B.
1NH ₄ OH . . .	12.3, T.; 12.4, B.	15.2	12.3, T.	1.3, B.	14.3, T.B.	8.4, T.; 5.3, B.	13.5, B.	12.7
$\frac{1}{2}$ Ca(OH) ₂ . . .	14.0, B.	18.4 †	13.9, B.	3.2	15.6, T.	9.3, † T.; 9.8, † B.	—	—
$\frac{1}{2}$ Sr(OH) ₂ . . .	13.8, T.	17.8 †	13.9, B.	3.15	15.4, T.	10.4, † T.B.	—	—
$\frac{1}{2}$ Ba(OH) ₂ . . .	13.9, B.	16.1	14.1, T.; 13.9, B.	3.15	18.4, B.T.	11.0, † T.B.	—	—
$\frac{1}{2}$ Mg(OH) ₂ . . .	13.8, B.	15.2	13.8, T.	1.5	15.3, B.T.	8.95, † B.	—	—
$\frac{1}{2}$ Cu(OH) ₂ . . .	7.5, T.	10.1	7.6	—	9.2	—	—	—

* 3NaOH gives 34.0 $\times 10^3$, T.

† Base in solid state.

‡ 1H₂SO₄.§ 1H₂CO₃.

HEATS OF COMBUSTION

HEATS OF COMBUSTION AND FORMATION OF CARBON COMPOUNDS, COAL, ETC.

Molecular heats of formation (H.F.) of organic compounds are deduced from their heats of combustion (H.C.), by subtracting the latter from the heat generated on burning the carbon and hydrogen contained in the compound. Experimental errors in the H.C. thus become magnified in the H.F. Heats of combustion determined by Thomsen are for the vapour of the compound at 18° C.; for the liquid the H.C. and H.F. would be greater by the latent heat of evaporation. Thomsen assumes H.F. of CO₂ from amorphous C as = 96,960 cal.; of water as 68,360 cal. per gm. molecule. For H.F. of inorganic compounds, see p. 62.

The H.C. and H.F. of carbon compounds is an **additive property** (see Thomsen's "Thermochemistry"). Berthelot's bomb calorimeter has been of considerable importance in the modern experimental side of the subject.

Unit—the gram calorie (at 15° to 20°) per gram molecule.

Example.—16 gms. of methane, CH₄, give out 212,000 gram calories of heat when burnt at **constant pressure**, to water and CO₂ at 18° C.

(T., Thomsen, "Thermochemistry;" B., Berthelot.)

Compound.	H.C.	H.F.	Compound.	H.C.	H.F.
	× 10 ³	× 10 ³		× 10 ³	× 10 ³
Methane, CH ₄ . . .	212, T.	21·7	Me. acetate, C ₃ H ₆ O ₂ . . .	399, T.	96·7
	213, B.		Carb. bisulphide, CS ₂ . . .	265, T.	-26
Ethane, C ₂ H ₆ . . .	370, T.	28·6	Methylamine, CH ₃ N . . .	258, T.	9·5
	372, B.		Dimethylamine, C ₂ H ₇ N . . .	420, T.	12·7
Propane, C ₃ H ₈ . . .	529, T.	35·1	Aniline, C ₆ H ₅ N . . .	838, T.	-17·4
	310, T.		Pyridine, C ₅ H ₅ N . . .	675, T.	-19·4
Acetylene, C ₂ H ₂ . . .	314	-47·8	Sugar, C ₁₂ H ₂₂ O ₁₁ . . .	1364	—
Ethylene, C ₂ H ₄ . . .	333, T.	-2·7	Illuminating gas per	5600 to	—
Benzene, C ₆ H ₆ . . .	799, T.	-12·5	cub. metre . . .	6500	—
Naphthalene, C ₁₀ H ₈ . . .	1239	—	Coal (anthracite) . . .	7·6 to	per gm.
				8·4	
Toluene, C ₇ H ₈ . . .	956, T.	-3·5	Coal (brown) . . .	4·7	" "
Me. alcohol, CH ₃ O . . .	182, T.	51·4	Coke . . .	6·9	" "
Me. chloride, CH ₃ Cl . . .	177, T.	22·6	Paraffin oil . . .	9·8	" "
Chloroform, CHCl ₃ . . .	107, T.	24·1	Wood . . .	{3·9 to}	" "
Et. alcohol, C ₂ H ₅ O . . .	340, T.	58·5		4·4	" "
Et. ether, C ₄ H ₁₀ O . . .	660, T.	70	Albumens—		
Et. chloride, C ₂ H ₅ Cl . . .	334, T.	30·7	Casein . . .	5·86	" "
Acetic aldehyde, C ₂ H ₄ O . . .	282, T.	48·7	Flesh . . .	5·66	" "
Formic acid, CH ₂ O ₂ . . .	69·4, T.	95·9	White of egg . . .	5·67	" "
Acetic acid, C ₂ H ₄ O ₂ . . .	225, T.	105·3	Yolk of egg . . .	8·12	" "
Propionic acid, C ₃ H ₆ O ₂ . . .	387, T.	109·4	Hæmoglobin . . .	5·9	" "
Me. formate, C ₂ H ₄ O ₂ . . .	241, T.	89·4			

MOLECULAR HEAT OF DILUTION

The heat set free or absorbed on diluting a gram molecule of liquid with water is the molecular heat of dilution: thus on diluting HCl to (HCl, 300 H₂O), 17,300 calories per 36·5 grams of HCl are set free; diluting 2NaCl, nH₂O (n = 20) to (2NaCl, 100H₂O) absorbs 1060 cal. per 2 × 58·65 gm. of NaCl. **Unit**—the gram calorie (at 15° to 20°) per gram molecule. (See Thomsen, "Thermochemistry.")

HCl n = 0	HNO ₃ n = 0	H ₂ SO ₄ n = 0	NaHO n = 3	NH ₃ * n = 1	2NaCl n = 20	2NaNO ₃ n = 12	Na ₂ SO ₄ n = 50	ZnCl ₂ n = 5	Zn(NO ₃) ₂ n = 10
H ₂ O × 10 ³	H ₂ O × 10 ³	H ₂ O × 10 ³	H ₂ O × 10 ³	H ₂ O × 10 ³	H ₂ O × 10 ³	H ₂ O × 10 ³	H ₂ O × 10 ³	H ₂ O × 10 ³	H ₂ O × 10 ³
1 5·37	1 3·28	1 6·38	5 2·13	1 1·26	100 -1·06	50 -2·26	100 -6·65	10 1·85	15 1·91
2 11·36	5 6·6	5 13·1	7 2·9	3 3·85	200 -1·31	100 -3·26	200 -1·13	20 3·15	20 1·15
5 14·96	10 7·32	49 16·7	9 3·1	5 8	400 -1·41	200 -3·86	400 -1·38	50 5·32	50 1·20
50 17·1	20 7·46	199 17·1	25 3·26	9 5	—	400 -4·19	800 -1·48	100 6·81	100 1·11
300 17·3	320 7·49	1600 17·9	200 2·94	110 0·00	—	—	—	400 8·02	200 1·07

* Heat developed on diluting NH₃.nH₂O to NH₃.200H₂O (Berthelot).

ENERGY AND WAVE-LENGTH OF FULL RADIATION

The radiation from a full or black body radiator depends both in quality and quantity upon the temperature. The total energy radiated (of all wave-lengths), from unit area in unit time, is given by *Stefan's law*, $E = K\theta^4$, where K is Stefan's constant and θ is the absolute temperature (see Optical Pyrometry, p. 47, and below).

The dependence of the quality on the temperature is expressed by *Wien's displacement law*, $\lambda_m\theta = \text{const.}$, where λ_m is the length of the particular waves which have maximum emissive power. Thus the emissive power E_m of the waves of length λ_m , varies as the 5th power of the temperature (absolute): $E_m\theta^{-5} = \text{const.}$

The emissive power of some particular wave-length λ is expressed accurately by

$$E_\lambda = C\lambda^{-5}/(e^{a/\lambda\theta} - 1) \quad \dots \dots \text{Planck's formula}$$

where $C = 353 \text{ erg. cm.}^2 \text{ sec.}^{-1}$, $a = 1.431 \text{ cm. deg.}$, and e is the base of Napierian logs.

At low temperatures or for short wave-lengths ($\lambda\theta < 3 \text{ cm. deg.}$) Planck's formula becomes (to '8 % at least)—

$$E_\lambda = C\lambda^{-5}e^{-a/\lambda\theta} \quad \dots \text{Wien's formula (see p. 47)}$$

For long waves and high temperatures ($\lambda\theta > 730 \text{ cm. deg.}$), we have (to 1 % at least)—

$$E_\lambda = C\lambda^{-4}\theta e^{-a/\lambda\theta} \quad \dots \dots \text{Rayleigh's formula}$$

(See Preston's "Heat," 2nd edit.; Kayser's "Spectroscopic," II.; Lorentz's "Theory of Electrons," 1910.)

WIEN'S DISPLACEMENT LAW		STEFAN'S LAW	
$\lambda_m\theta = \text{const.} = A.$ (See above). λ is measured in cms.		Total radiation from a full radiator $= K\theta^4$ (see above). K is in $\text{erg cm.}^{-2} \text{ sec.}^{-1} \text{ deg.}^{-4}$. $K = 5.72 \times 10^{-5}$ (Millikan, 1917).	
A	Observer.	K	Observer.
2940	Lummer and Pringsheim, 1899	5.45×10^{-5}	Kurlbaum, 1912
2888	Paschen and Wanner, <i>B. B.</i> , 1899	5.18	{ Lummer and Pringsheim, <i>A. d. P.</i> , 1901
2902	Wanner, 1900	5.67	
2940	Paschen, <i>A. d. P.</i> , 1901	5.89	Shakespeare, 1911
2890	Rubens and Kurlbaum, <i>A. d. P.</i> , 1901		Keene, 1912

A. d. P., *Ann. der Phys.*; *B. B.*, *Berlin Ber.*; *C. R.*, *Compt. Rend.*

SOLAR CONSTANT AND TEMPERATURE OF SUN

The solar constant S is the energy received from the sun by the earth (at its mean distance) per sq. cm. in unit time, corrected for the loss by absorption in the earth's atmosphere.

The determination of the absorption loss is difficult; it is best derived from simultaneous observations at high and low stations.

Langley and Abbot ("Smithsonian Reports," 1903 *et seq.*) give the following relation between atmospheric absorption and wave-length:—

Wave-length (\AA .U. = 10^{-8} cm.)	4000	6000	8000	10,000	12,000
Fraction transmitted	.49	.74	.85	.89	.91

If R is the energy radiated in unit time from a sq. cm. of the sun's surface, then

$$R = \left\{ \frac{\text{earth's solar distance}}{\text{sun's radius}} \right\}^2 \times S = \left\{ \frac{9.28 \times 10^7}{4.33 \times 10^5} \right\}^2 \times S = 46,000S$$

Assuming the sun to be a full or black body radiator, its "effective" absolute temperature θ may be deduced either from (1) Stefan's law, $R = K(\theta^4 - T^4)$, where K is Stefan's constant (see above) and T is the earth's absolute temperature, or (2) Wien's displacement law, $\theta\lambda_m = \text{const.}$ (see above).

Langley and Abbot (ref. above) find the distribution of the energy of solar radiation among the different wave-lengths (λ) to be as follows:—

Wave-length (\AA .U.)	4000	4500	5000	5500	6000	7000	8000	10,000	12,000	14,500	21,000
Relative energy, E	15.2	18.4	19	16	14	11	8.8	5.4	3.2	2	.6

λ for $E_{\text{max.}} = 4900 \times 10^{-8} \text{ cm.}$ Taking Wien's displacement law to be $\theta\lambda_{\text{max.}} = .29$, and assuming the sun to be a full radiator, its temperature $\theta = 5920^\circ$ absolute.

SOLAR CONSTANT

SOLAR CONSTANT AND TEMPERATURE OF THE SUN (*contd.*)

The values of S below are expressed in both (1) calories per min. per cm^2 , and (2) watts per cm^2 (1 calorie per sec. = 4.18 watts). The sun's mean temp. θ is in degrees C. absolute. Abbot and Fowle find the solar constant varies by about 8 % (See Poynting and Thomson's "Heat;" Chree, *Nature*, 82, 2090; Report (1910) of the International Union for Solar Research; and "Smithsonian Reports.")

Solar Const.		Sun's Temp.	Account.	Observer.
cal. min. ⁻¹ cm. ⁻²	watts cm. ⁻²			
—	—	Abs. 5770°	Comparison with const. temp. Atmos. absorp. taken as 29 %	Wilson, 1902
—	—	5920	Using Wien's displacement law (above)	Langley & Abbot, '03
2.25	.154	7060	Gorner Grat, Switzerland	Scheiner, 1908
—	—	5610	Natl. Phys. Lab., England. Atmos. absorp. taken as 29 %	Harker & Blackie, '08
2.38	.166	5630	Mt. Blanc. Comparison with const. temp. Atmos. absorp., 9 % with zenith sun	(Féry & Millochau
—	—	5360		Féry, 1909
—	—	5630		Millochau, 1909
2.1	.146	5970†	Mt. Blanc. Atmos. absorp., 3.4 % Washington (sea-level) and Mt. Wilson (6000 ft.)	Abbot & Fowle, '09
2.1	.146	5970†	Review of previous work	Bellia, 1910
1.925*	.134	5840†	Mt. Wilson (6000 ft.) and Mt. Whitney (14,500 ft.)	Abbot, 1910

* Mean value for period 1904-9 (*Nature*, 1911).

† Calculated from S , taking Stefan's const. as 5.3×10^{-12} watts cm^{-2} sec.⁻¹ deg.⁻⁴.

THE CRYOSCOPIC CONSTANT

The cryoscopic constant, K , would be the depression of the freezing-point of a solvent when the molecular weight in grams of any substance (which does not dissociate or associate) is dissolved in 100 grams of the solvent, supposing the laws for dilute solutions held for such a concentration (Raoult, 1882). Van't Hoff (1887) showed that $K = R\theta^2/(100L)$, where R = the gas constant (see p. 5), θ the absolute freezing-point of the solvent, L its latent heat of fusion in ergs. **Example.**—For 1 gram-molecule of solute in 100 gms. of water—

$$K = 8.315 \times 10^7 \times (273.1)^2 / (79.67 \times 4.184 \times 10^9) = 18.60$$

(See Whetham's "Theory of Solution," p. 149.)

Solvent.	M. pt.	Lat. ht. (cals.)	K		Solvent.	M. pt.	Lat. ht. (cals.)	K	
			Calcd.	Obsd.				Calcd.	Obsd.
Water . .	0° C.	79.6	18.6	18.58, G. 18.52 *	Benzene .	5° C.	29.1, P.W. 30.1, F.	53.3 51.6	49, R. 51.2, P.
H ₂ SO ₄ , H ₂ O	8.4	31.7, B.	50	.48, L.	Formic acid	8	57.4, Pe.	27.5	28, R.
SbCl ₃ . .	73.2	13.4, T.	174	184, T.	Phenol . .	40	24.9, P.W.	78.6	72.7, E.
Acetic acid	17	43.7, Pe.	38.5	39, R.	p. Xylol .	16	39.3, C.	42.5	43, P.M.
Aniline . .	- 6	—	—	58.7, A.R.					

* Mean of six observers; A.R., Ampola and Rimatori, 1897; B., Berthelot; C., Colson; E., Eykman, 1889; F., Fischer; G., Griffiths (who used 0.0005 to 0.02 normal sugar solutions); L., Lespieau, 1894; P., Paternò, 1889; Pe., Pettersson; P.M., Paternò and Montemartini, 1894; P.W., Pettersson and Widman; R., Raoult; T., Tolloczko, 1899.

VELOCITY OF SOUND

The velocity of sound (longitudinal waves) in a body, $V = \sqrt{E/\rho}$, E being the elasticity, and ρ the density. In gases and liquids E is the adiabatic volume elasticity; in isotropic solid rods or pipes E is Young's Modulus. For gases $V = \sqrt{\gamma P/\rho}$, P being the pressure, and γ the ratio of the specific heat of the gas at constant pressure to that at constant volume. For values of γ , see p. 58.

For moderate temperature variations, the velocity of sound in gases is given by $V_t = V_0(1 + \frac{1}{2}at) = V_0 + 61t$ in cms. per sec. for dry air ($a = .00367$).

The velocity of sound decreases with decreasing intensity down to the normal value. In gases in tubes the velocity increases with the diameter up to a limiting value for free space. The values below are for free space. Barton's "Sound" and Poynting and Thomson's "Sound" may be consulted. [1 foot = 30.48 cms.]

Substance.	Temp.	Velocity.	Observer.
Gases—			
Air (dry)	0° C	$(3.3133) \times 10^4$	Calcd. ($\gamma = 1.401$)
"	0	3.3136 "	Violle, 1900
"	0	3.3132 "	Stevens, 1900
"	0	3.3129 "	Hebb, 1905
"	0	3.3192 *	Thiesen, 1908 ‡
"	- 45.6	3.056 "	Greely, 1890
"	- 182.4	1.815 "	Cook, 1906
"	100	3.865 "	Stevens, 1900
"	500	5.53 "	"
"	1000	7.0 "	"
" (Krakatoa wave)	—	3.21 "	1883
" Sound-waves from sparks	0	3.50-4.45 ", †	Töpler, 1908
Hydrogen	0	12.86 "	Zoch, 1866
Oxygen	0	3.172 "	Dulong, 1829
"	- 184.7	1.737 "	Cook, 1906
Nitrous oxide, N_2O	0	2.60 "	Wullner, 1878
Ammonia, NH_3 . .	0	4.16 "	"
Carbon monoxide . .	0	3.371 "	"
Carbon dioxide . .	10-24	2.573 "	Low, 1894
Coal-gas	0	4.9-5.15 "	—
Sulphur dioxide . .	0	2.09 "	Masson, 1857
Water-vapour . . .	0	4.0 "	"
" (satd.)	110	4.13 "	Treitz, 1903
Liquids—			
Water	8.1	14.35×10^4	Colladon & Sturm, 1827
"	4	13.99 "	Martini, 1888
"	25	14.57 "	"
" (sea) Explosion	waves 18	17.3-20.1 ", †	Threlfall & Adair, 1889
Alcohol (abs.), C_2H_5O	8.4	12.6 "	Martini, 1888
Ether, $(C_2H_5)_2O$. .	0	11.4 "	"
Turpentine, $C_{10}H_{18}$	3.5	13.7 "	"

* Free from CO_2 . † The range of speeds is given by varying intensities. ‡ Reichsanstalt.

The values for metals are due to Wertheim, 1849; Masson, 1857; and Gerossa, 1888.

Solid.	Velocity cms./sec.	Solid.	Velocity cms./sec.	Solid.	Velocity cms./sec.
Aluminium. . .	51.0×10^4	Lead	12.3×10^4	Brass	$c. 36.5 \times 10^4$
Cadmium . . .	23.1 "	Nickel	49.7 "	Deal (along	49-50 "
Cobalt	47.2 "	Platinum . . .	26.8 "	grain)	
Copper	39.7 "	Silver	26.4 "	Fir	42-53 "
Gold	20.8 "	Tin	24.9 "	Mahogany . .	41-46 "
Iron (wrought)	49-51 "	Zinc	36.8 "	Oak	40-44 "
" (cast)	c. 43 "	Glass (soda)	50-53 "	Pine	c. 33 "
Steel	47-52 "	" (flint) . . .	c. 40 "	Indiarubber .	5-7 "

VELOCITY (IN AIR) AND PRESSURE Koch (1907).			SENSITIVENESS OF EAR TO PITCH Rayleigh (1907).		ORGAN PIPES End Correction.	
Press. in atmos.	Relative Velocity of Sound.		Frequency.	Condensation for same audibility.	<p>For a pipe with a flange at the open end, the antinode is situated .82 (radius of pipe) beyond end. With no flange, the end-correction is .57 (radius). (See Lamb's "Sound," [1910.]</p> <p>Wave-length. L = length of pipe.</p> <p>Closed pipe . . . 4L, $\frac{4L}{3}$, $\frac{4L}{5}$, etc. Open pipe . . . 2L, $\frac{2L}{2}$, $\frac{2L}{3}$, etc.</p>	
	0° C.	-79.3° C.				
1	1.000	.842	512	1		
.25	1.008	.831	256	1.6		
50	1.022	.830	128	3.2		
100	1.064	.885	85	6.4		
150	1.132	1.047				
200	1.220	1.239				

TRANSVERSE VIBRATIONS OF RODS				THE EAR	
L, length; K, radius of gyration of cross-section; E, Young's Modulus; ρ, density.					
	No. of Nodes.	Distance of Nodes from one end.	Frequency $\propto \frac{K}{L^2} \sqrt{\frac{E}{\rho}}$		
Both ends free	2	.224 L; .776 L	1	Shortest time perceivable by ear (Hill, 1908) '007 sec.	
	3	.132 L; .5 L; .868 L	2.76	Amplitude of faintest audible sound (Rayleigh, 1877) 10.4×10^{-8} cm.	
	4	{ .094 L; .356 L } { .644 L; .906 L }	5.40	Ditto (Shaw, 1904) 1.4×10^{-8} cm.	
One end fixed	0	—	1	Pressure variation to which normal ear can respond (Abraham, 1907) 6.4×10^{-7} mm. mercury.	
	1	.226 L	6.27	Lower limit of audition in vibns./sec. About 30.	
	2	.132 L; .5 L	17.5	Upper limit of audition in vibns./sec. 24,000 to 41,000.	
	3	.094 L; .356 L; .644 L	34.4	Extreme range of ear Musically available c. 11 octaves. c. 7 "	
Temp. correction of Frequency (n) of a Tuning-fork. (M'Leod and Clarke, 1880, and König) $n_t = n_0(1 - .00011t)$				Highest pitch in piano 3520	
The pressure exerted by Sound waves has been measured directly up to .24 dyne/cm ² . (Altberg, 1903)				Highest pitch in orchestra (piccolo d'). 4752	
				Lowest pitch in largest organs (64-foot pipe) 8	

FREQUENCY RATIOS OF MUSICAL SCALE									
	C Doh	D Ray	E Me	F Fah	G Soh	A Lah	B Te	c Doh	
Natural scale	1	$\frac{9}{8}$	$\frac{10}{9}$	$\frac{16}{15}$	$\frac{9}{8}$	$\frac{10}{9}$	$\frac{8}{7}$	$\frac{16}{15}$	2
	24	27	30	32	36	40	45	48	
	1.000	1.125	1.250	1.333	1.500	1.667	1.875	2.000	
Equally tempered scale	1.000	1.122	1.260	1.335	1.498	1.682	1.888	2.000	
Standard forks (König) (marked c' = 512 and so on)	c'	d'	e'	f'	g'	a'	b'	c''	
	256	288	320	341.3	384	426.7	480	512	

The French Standard, "Diapason Normal" of 1859 (which adopts a fork having $c'' = 522$ at 20° C.) is coming into general adoption for organs and pianos in England, the Continent, and America, as the result of a makers' conference in 1899. Other scales in vogue are Concert Pitch ($c'' = 546$), Society of Arts ($c'' = 528$), Tonic Sol-fa ($c'' = 507$), Philharmonic ($c'' = 540$). (The "middle" c of the piano is c'.)

VELOCITY OF LIGHT IN VACUO

Mean value *in vacuo* = 2.9986×10^{10} cm./sec. = 186,326 miles/sec. For values of v , the ratio between the E.M. and E.S. units, see below.

cm./sec.	Method.	Observer.	cm./sec.	Method.	Observer.
$\times 10^{10}$			$\times 10^{10}$		
3'07	Eclipse of one of Jupiter's moons	Römer, 1676	2'999	Rotating mirror	Michelson, 1879
2'998		„ corrected	3'014	Toothed wheel	Young & Forbes, '81
3'153	Toothed wheel	Fizeau, 1849	2'9985	Rotating mirror	Michelson, 1882
2'986	Rotating mirror	Foucault, 1862	2'9986	„	Newcomb, 1882
3'004	Toothed wheel	Cornu, 1878	2'9986	Toothed wheel	Perrotin, 1900

VELOCITY OF LIGHT IN LIQUIDS

Liquid.	Vel. in vacuo Vel. in liquid	Refractive index for Na D line.	Method.	Observer.
Water . .	1'330	1'333/20°	Rotating mirror	Michelson, 1883
CS ₂ . . .	1'758	1'627/20°	„ „	„ „

VELOCITY OF HERTZIAN WAVES

(See Blondlot and Gutton, *Rep. Cong. Phys.*, Paris, 1900.)

cm./sec.	Observer.	cm./sec.	Observer.	cm./sec.	Observer.
$\times 10^{10}$		$\times 10^{10}$		$\times 10^{10}$	
2'989	Blondlot	3'003	Trowbridge	2'989	Saunders
2'991	McClean		and Duane	2'991	Mean

RATIO OF ELECTROMAGNETIC TO ELECTROSTATIC UNIT OF CHARGE

This ratio " v " is a pure number, and is numerically equal to $\sqrt{\mu k}$, i.e. on Maxwell's theory, to the velocity of electric disturbances, such as light and Hertzian waves, through a medium whose magnetic permeability is μ and specific inductive capacity k . (See pp. 7 and 84.) For the velocity of light, see above.

Most observers have used a "capacity method" of determining v . (See Gray, "Absolute Measurements"; and Rosa, *Bull. Bureau of Standards*, 1907.)

v	Observer.	v	Observer.	v	Observer.
$\times 10^{10}$		$\times 10^{10}$		$\times 10^{10}$	
2'963	J. J. Thomson, 1883	2'997	Thomson and Searle, 1890	3'001	Hurmuzescu, '96
2'982	Rowland, 1889	3'009	Pellat, 1891	2'997	Perot and Fabry
3'000	Rosa, 1889	2'993	Abraham, 1892	2'997	Rosa & Dorsey, 1907

PHOTOMETRIC STANDARDS

The Geneva Congress of 1896 proposed a set of units for measuring (1) luminous intensity, (2) flux (the "lumen"), (3) illumination (the "lux"), (4) brightness, and (5) quantity of light (see *Electrician*, July 14, 1911). The British unit of intensity is the "candle." The **mean spherical candlepower** of a light is the mean of the intensities measured in all directions from the light. The **mean horizontal candlepower** is the mean of all the intensities in a horizontal plane through the lamp.

The **British "candle"** is a spermaceti candle, $\frac{7}{8}$ inch in diameter (6 to the lb.) which burns at the rate of 120 grains per hour. This is, however, found to be an unsatisfactory standard, and in modern photometry the British unit is taken as being one-tenth part of the light given out by the Harcourt 10 candlepower Pentane lamp, burning at a pressure of 760 mms. mercury in an atmosphere containing 8 parts in 1000 by volume of water-vapour as measured by a ventilated hygrometer. The candlepower of this lamp

$$= 10 + \cdot 066(8 - w) - \cdot 008(760 - H)$$

where w is the number of parts in 1000 (by vol.) of water-vapour in air at a barometric pressure of H mms. of mercury.

The **United States "candle"** prior to April 1, 1909, was 1·6% greater than the British:

The **French unit** is the Bougie decimale, which is the 20th part of the light given out by a sq. cm. of platinum at its solidifying point. This is a difficult unit to reproduce, and the Carcel lamp burning colza oil is used in practice. The Carcel unit is taken (with some uncertainty) as 4% less than the Bougie decimale.

The **German unit** is the light given out by the Hefner lamp (which burns amyl acetate), burning at a pressure of 760 mms. mercury in an atmosphere containing 8·8 parts in 1000 (by vol.) of water-vapour as measured by a ventilated hygrometer.

The National Physical Laboratory, the Bureau of Standards of America, and the Laboratoire Central d'Electricité of Paris have come to an agreement which involves the reduction of the old value of the American candle by 1·6%. They agree in future to employ as a common unit the proposed **International candle** = 1 British Pentane candle = 1 American candle = 1 French Bougie decimale = 10/9 German Hefner unit = 104 Carcel unit (see Paterson, *Phil. Mag.*, 1909).

EFFICIENCIES OF VARIOUS LIGHTS

It has become customary to express efficiencies (or rather inefficiencies) in watts per candle. The value of a luminous efficiency cannot be properly appreciated without a knowledge of the distribution of the intensity. Estimates of the proportion of light energy to the total energy vary widely. S. P. Thompson ("Manufacture of Light") quotes from 1 part in 7000 for a gas flame to 1% for the most efficient lights.

The usual accepted "efficiencies" are given below in watts per mean spherical candlepower. They must only be regarded as approximate (see Solomon, "Electric Lamps," 1908).

Light.	Efficiency.	Light.	Efficiency.
Bat's-wing gas flame	c. 100	Tantalum lamps	1'7-2'1
Paraffin lamps	c. 50	Tungsten (osram, etc.) lamps	1'3
Welsbach mantle, etc.	c. 15	Open arc lamps	1'1-1'4
High-pressure gas	c. 8	Enclosed arc lamps	2'3
Carbon filament lamps	3'5-4'5	Yellow flame arc lamps	'4
Metallized carbon filament lamps	2'8	Mercury vapour lamps	'3-'4
Nernst lamps	2'1-2'4		

In high-grade standard photometry the Lummer Brodhun photometer head is usually employed. A unit of light may be maintained and reproduced with an accuracy of the order of $\frac{1}{10}$ %, by means of sets of properly seasoned glow lamps.

The candlepower of a carbon glow lamp varies as the 6th power (approx.) of the voltage; of a metallic filament lamp, as the 3/6th power.

A candle is visible at about a mile on a clear dark night. The energy in the luminous radiation from a standard candle is about 5×10^5 ergs/sec. (Rayleigh, "Collected Papers"), whence the energy falling on 1 sq. cm. at a distance of 1 metre would be 4 ergs per sec. Angström (1902) gets values about double these.

71
GASEOUS REFRACTIVE INDICES

GASEOUS REFRACTIVE INDICES AND DISPERSIONS

Dispersion.—Cauchy's equation is $\mu - 1 = A(1 + B/\lambda^2)$, where μ is the refractive index for the wave-length λ ; A and B are constants. B is the coefficient of dispersion.

The **refractivity** $(\mu - 1) = A$, when $\lambda = \infty$. The values of A and B are for wave-lengths measured in cms. The refractive indices are mostly for the sodium D line ($\lambda = 5893 \times 10^{-8}$ cm.). The values of μ are reduced to a standard density at 0° and 760 mms. by assuming that $(\mu - 1)/\rho$ is a constant for each gas, ρ being the density. Cauchy's formula is in general inadequate over large dispersions. (See Cuthbertson, *Science Progress*, 1908; and *Proc. & Trans. Roy. Soc. for 1905 et seq.*)

Gas or Vapour.	Refractive Index μ for Na D line.	Cauchy's Constants.		Observer.
		A.	B.	
Air . . .	1'0002918	28.71×10^{-5}	5.67×10^{-11}	Scheel (Reichsanstalt), 1907.
Hydrogen . .	1'0001384	13.58 "	7.52 "	"
Helium . . .	1'0000350	3.48 "	2.3 "	Burton; Cuthbertson & Metcalfe, 1907
Neon . . .	1'0000671	6.66 "	2.4 "	C. & M. Cuthbertson, 1909
Argon . . .	1'0002837	27.92 "	5.6 "	Burton, 1907
Krypton . .	1'0004273	41.89 "	6.97 "	C. & M. Cuthbertson, 1908
Xenon . . .	1'000702	68.23 "	10.14 "	"
Fluorine . .	1'000195	—	—	Cuthbertson & Prideaux, 1906
Chlorine . .	1'000768	—	—	Mascart, 1878
Bromine . .	1'001125	—	—	"
Iodine . . .	1'00192 †	—	—	Hurion, 1877
Oxygen . . .	1'000272	26.63 "	5.07 "	Rentschler, 1908
Sulphur . .	1'001111	104.6 "	21.2 "	Cuthbertson & Metcalfe, 1908
Selenium . .	1'001565	—	—	"
Tellurium . .	1'002495	—	—	"
Nitrogen . .	1'000297	29.06 "	7.7 "	Scheel (Reichsanstalt), 1907
Phosphorus .	1'001212	116.2 "	15.3 "	Cuthbertson & Metcalfe, 1908
Arsenic . . .	1'001552	—	—	"
Zinc . . .	1'002050	—	—	"
Cadmium . .	1'002675	—	—	"
Mercury . .	1'000933	87.8 "	22.65 "	"

Gas or Vapour.	Refractive Index μ for Na D line.	Observer.	Gas or Vapour.	Refractive Index μ for Na D line.	Observer.
Water-vapour . .	1'000257	Mascart, '78	Tellurium tetra-chloride . . .	1'002600	P. & M.
" " . . .	1'000250	Lorenz, '74	Phosph. hydrogen	1'000786*	Dulong, '26
Ammonia . . .	1'000377	Mascart, '78	Phosphorus tri-chloride . . .	1'001730	Mascart, '78
" " . . .	1'000373	Lorenz, '74	Methane, CH ₄ . .	1'000441	" "
Nitrous oxide . .	1'000515	Mascart, '78	Pentane, C ₅ H ₁₂ . .	1'001701	" "
Nitric oxide . .	1'000297	" "	Acetylene, C ₂ H ₂ . .	1'000606	" "
Hydrochloric acid	1'000444	" "	Ethylene, C ₂ H ₄ . .	1'000719	" "
Hydrobromic acid	1'000570	" "	" " . . .	1'000674	Prytz, '80
Hydriodic acid .	1'000906	Hurion, '77	Benzene, C ₆ H ₆ . .	1'001812	Mascart, '78
Carbon monoxide	1'000334	Mascart, '78	" " . . .	1'001765	Prytz, '91
" dioxide . . .	1'0004498	Perreau, '96	Methyl fluoride . .	1'000449	Cuthbertson
" bisulphide .	1'001476	Mascart, '78	" chloride . . .	1'000865	Mascart, '78
Sulph. hydrogen	1'000641*	Dulong, '26	" alcohol . . .	1'000552	Prytz, '80
" " . . .	1'000619	Mascart, '78	" " . . .	1'000619	Mascart, '78
Sulphur dioxide .	1'000660	Walker, '03	Chloroform, CHCl ₃	1'001455	" "
" trioxide . .	1'000737	C. & M., '08	Carbon tetra-chloride . . .	1'001768	" "
" hexafluoride	1'000783	"			
Selenium " . .	1'000895	"			
Tellurium " . .	1'000991	"			

* White light. † Violet light. $\mu = 1.00205$ for red light. Iodine shows anomalous dispersion. C. & M., Cuthbertson & Metcalfe; P. & M., Prideaux & Metcalfe.

REFRACTIVE INDICES

REFRACTIVE INDICES

Refractive indices, μ , (against air) at 15° C. for various wave-lengths.

The **temperature coefficient** given below is the change of refractive index per 1° C. rise of temperature for the case of the sodium D line.

The refractive indices are due chiefly to Gifford (*Proc. Roy. Soc.*, 1902, 1904, 1910); Rubens and Paschen (for the infra-red) and Martens (1902). The two Jena glasses are selected as typical. Other glasses are dealt with on p. 74.

Wave-length in Å.U. (10^{-8} cm.).	Calcsp. 18°.		Jena glass.		Flu- orite, CaF ₂ , 18°.	Quartz, 18°.		Fused silica.	Rock salt, 18°.	Syl- vin, KCl 18°.	Water at 20°.
	ord. ray.	ext. ray.	Crown*	flint.†		ord. ray.	ext. ray.				
Infra-red.	1'	1'	1'	1'	1'	1'	1'	1'	1'	1'	1'
223,000	—	—	—	—	—	—	—	—	3403	3712½	—
94,290	—	—	—	—	3161	—	—	—	4983	4587	—
42,000	—	—	—	—	4078	4569	—	—	5213	4720	—
21,720	6210	4746	4946	6153	4230	5180	5261	—	5262	4750	—
12,560	6388	4782	5042	6268	4275	5316	5402	—	5297	4778	3210
Visible.											
Li, (r) 6708	6537	4843	5140	6434	4323	5415	5505	4561	5400	4866	3308
H, (C) 6563	6514	4846	5145	6414	4325	5419	5509	4564	5407	4872	3311
Cd, (r) 6438	6550	4847	5149	6453	4327	5423	5514	4568	5412	4877	3314
Na, (D) 5893	6584	4864	5170	6499	4339	5443	5534	4585	5443	4904	3330
Hg, (g) 5461	6616	4879	5191	6546	4350	5462	5553	4602	5475	4931	3345
Cd, (g) 5086	6653	4895	5213	6598	4362	5482	5575	4619	5509	4961	3360
H, (F) 4861	6678	4907	5230	6637	4371	5497	5590	4632	5534	4983	3371
Cd, (b) 4800	6686	4911	5235	6648	4369	5501	5594	4636	5541	4990	3374
Hg, (v) 4047	6813	4969	5318	6852	4415	5572	5667	4697	5665	5097	3428
Ultra-violet.											
Sn 3034	7196	5136	5552	—	4534	5770	5872	4869	6085	5440	3581
Cd 2144	8459	5600	—	—	4846	6305	6427	5339	7322	6618	4032
Al 1852	—	—	—	—	5099	6759	6901	5743	8933	8270	—
Temp. co- efficient (D)	+ '005	+ '0414	- '001	+ '003	- '041	- '005	- '006	- '003	- '04	- '04	- '048

* Light barium crown. † Dense silicate flint. ‡ $\mu = 1.3692$ for $\lambda = 225,000$.

REFRACTIVE INDICES

Refractive indices μ_D (against air) at 15° C. for sodium D line ($\lambda = 5893 \times 10^{-8}$ cm.).

Substance.	μ_D	Substance.	μ_D	Substance.	μ_D
Solids.		Alcohol, ethyl . . .	1.362	Monobrom benzene	1.563
Alum (potash) . . .	1.456	" amyl . . .	1.41	" " naphtha- lene . . .	1.660
Cyanin	1.71	Aniline	1.590	Nitrobenzene . . .	1.553
Diamond	2.417	Benzene	1.504	Oil, cedar	1.516
Glass (see above and p. 74)	1.31	Bromoform	1.591	" cloves	1.532
Ice	1.60	Canada balsam . . .	1.53	" cinnamon	1.601
Mica . . . 1.56 to	1.76	Carb. bisulphide . .	1.632	" olive	1.46
Ruby	1.56	" tetrachloride . .	1.464	" paraffin	1.44
Sugar	1.63	Chloroform	1.449	Sulphuric acid . . .	1.43
Topaz	1.63	Ether, ethyl	1.354	Turpentine	1.47
Liquids.		Ethylene dibromide	1.540	Water (see above).	1.333
Alcohol, methyl . .	1.33	Glycerine	1.47		
		Methylene iodide . .	1.744		

DISPERSIVE POWERS

The dispersive power (ω) given below = $(\mu_C - \mu_F)/(\mu_D - 1)$, where μ_C , μ_D , μ_F are the refractive indices corresponding to the red (C) H line (6563), the yellow Na (D) line (5893), and the green-blue (F) hydrogen line (4862).

Substance.	ω	Substance.	ω	Substance.	ω
Solids.				Liquids.	
Calcite, ord. . . .	0204	Quartz, ord. . . .	0143	Carb. bisulphide . .	0545
„ ext. . . .	0125	„ ext. . . .	0146	Alcohol	0171
Fluorite	0105	Fused silica . . .	0147	Turpentine	0206
Glass (see p. 74)		Rock salt	0233	Water	0180
		Sylvin. . . .	0226		

SILVERING SOLUTION

Due to the late Dr. Common. Other recipes will be found in Baly's "Spectroscopy" (Longmans) and Woollatt's "Laboratory Arts" (Longmans).

Make up 10 % solutions of (1) pure nitrate of silver, AgNO_3 ; (2) pure caustic potash, KOH ; (3) loaf sugar; and (4) ammonia (90 % water, 10 % ammonia of sp. gr. 880). To the sugar soln. add $\frac{1}{2}$ % of pure nitric acid and 10 % of alcohol. The sugar soln. is very much improved by keeping. Make up also a 1 % soln. of AgNO_3 . Distilled water must be used for all the solns.

For silvering say a 12-in. mirror, take 400 c.c. of the AgNO_3 soln. and add strong ammonia until the brown precipitate first formed is nearly dissolved, then use the 10 % ammonia until the soln. is just clear. Add 200 c.c. of the KOH soln. A brown precipitate is again formed, which must be dissolved in ammonia exactly as before, the ammonia being added until the liquid is just clear. Now add the 1 % soln. of AgNO_3 until the liquid becomes a light brown colour about equal in density of colour to sherry. This colour is important, and can only be properly obtained by the use of the weak soln. Dilute the liquids to 1500 c.c. with distilled water.

The mirror should be thoroughly cleaned with acid and placed in a dish of distilled water.

All being ready, add 200 c.c. of the sugar soln. to 500 c.c. of water; add the mixture to the silver-potash soln., mix thoroughly, and pour them into a clean empty dish. Then lift the mirror out of its dish of distilled water and place it face downwards in this soln., taking care to exclude all air-bubbles.

The liquid will turn light brown, dark brown, and finally black. In four or five minutes, often sooner, a thin film of silver will commence to form on the mirror, and this will thicken until in about 20 minutes the whole liquid has acquired a yellowish-brown colour, with a thin film of metallic silver floating on the surface. Half an hour is the usual time taken in silvering, but this is shortened by using warmer liquids. About 18° C. is the best temperature.

Lift the mirror out, thoroughly wash with distilled water, and stand on its edge for say 12 hours in an inclined position until it is dry. The slight yellowish "bloom" can then be polished off by rubbing softly with a pad of chamois leather and cotton-wool. The subsequent polishing is done with a little dry well-washed rouge on the leather pad. The film should be opaque and brilliant, and with careful handling will be very little changed with long use.

Porcelain, glass, or earthenware dishes should be used.

If a very thick film is required, two silvering baths can be used, the article being left in the first bath for 15 minutes, then lifted out, rinsed with distilled water and at once immersed in the second bath, which should be ready in another dish. The film should not be allowed to dry during the operation of changing baths.

NOTE.—The silver-potash solution will not keep beyond a couple of hours. Any excess of this solution unused should have the silver precipitated at once with HCl . If the silver-potash is kept, say for 10 or 12 hours, a black powder collects on the surface. This powder, which is probably some form of fulminate of silver, is explosive, and may shatter the vessel.

GLASS

The **raw materials** for the manufacture of glass are (1) silica—usually as sand or felspar; (2) salts of the alkali metals— Na_2SO_4 , Na_2CO_3 , or K_2CO_3 ; (3) salts of bases other than alkalies—red lead, limestone or chalk, BaCO_3 or BaSO_4 , MgCO_3 , ZnO , MnO_2 , Al_2O_3 , As_2O_3 , etc. In general, glasses rich in silica and lime are hard, while glasses in which alkali, lead, or barium preponderate are soft. Hardness is, of course, also largely dependent on annealing. Ordinary “soft” (*i.e.* easily fusible) German glass is a soda-lime glass rather rich in alkali; “hard” (refractory) glass is a potash-lime glass rather rich in lime. Jena combustion tubing is a borosilicate containing some magnesia.

Thermometry Glasses.—Glasses which contain **both** soda and potash to any extent give a large temporary zero depression (see p. 45). Data concerning *Verre dur* (71% SiO_2 , 12% Na_2O , $\frac{1}{2}\%$ K_2O , 14% CaO , 2% Al_2O_3 and MgO), *Fena 16* (67% SiO_2 , 14% Na_2O , 7% CaO , 12% ZnO , Al_2O_3 and B_2O_3), *Fena 59* (72% SiO_2 , 12% B_2O_3 , 11% Na_2O , 5% Al_2O_3), *Kew glass* (44% SiO_2 , 34% PbO , 12% K_2O , 2% Na_2O , 2% CaO , MgO , etc.), will be found on p. 45.

Optical Glasses.—In building up achromatic lens systems a knowledge of the dispersive power (ω) of each glass employed is essential. This is defined as the ratio of the difference of the deviations (*i.e.* the dispersion) for any two colours to the deviation of some mean intermediate colour. ω thus depends on the colours selected; for visual work they are usually the red (C) line of hydrogen (wave-length $\lambda_C = 6563 \times 10^{-8}$ cm.), the yellow sodium (D) line ($\lambda_D = 5893$), and the green-blue (F) hydrogen line ($\lambda_F = 4862$). If μ_C , μ_D , μ_F are the corresponding refractive indices, $\omega = (\mu_C - \mu_F)/(\mu_D - 1)$ for the brightest part of the visible spectrum.

Flint glass—a term which survives from times when ground flints were extensively employed in making the best glass—now always implies a dense glass which contains lead and has a high refractive index and dispersive power.

Crown glass, originally designating only lime-silicate glasses, is now applied generally to glasses having a low dispersive power.

Jena Optical Glasses.—For ordinary flints and crowns ω and μ are roughly proportional, and this was true for all commercially available glasses prior to the advances initiated in 1881 by Abbé and Schott at Jena. They succeeded (*e.g.* by the addition of barium) in producing glasses which do not obey any such proportionality; *e.g.* the very valuable barium crown glasses (below) combine the high refractive index of a flint glass with the low dispersive power of a crown. Such glasses have brought about the excellent achromatism and flatness of field which now obtain in photographic lenses and large telescopic objectives. The introduction of boron into a glass lengthens the blue end of the spectrum relatively to the red; the addition of phosphorus, fluorine, potassium, or sodium has the opposite effect: such control over the dispersion has made the modern microscope possible.

Some typical examples of Jena glasses are subjoined. For a complete list, see the catalogue of Schott and Genossen, Jena. The simple phosphate and borate glasses have been withdrawn on account of their lack of durability. The borosilicate crowns are among the most durable and chemically resistant of all glasses. The U.V. glasses are markedly transparent to ultra-violet light as far as about $\lambda = 2880$.

See p. 72, and Zschimmer's “History of the Jena Glass Works,” Hovestadt's “Jena Glass,” and Rosenhain's “Glass Manufacture,” 1908 (with bibliography).

(After Zschimmer, *Zeit. Inst.*, 1908.)

Glass.	μ_D	ω (C,D,F)	Dens. grms. c.c.	Glass.	μ_D	ω (C,D,F)	Dens. grms. c.c.
Crowns—				Flints (contd.)—			
(Silicate) crown .	1.4782	.0152	2.23	U.V. flint 3492 . .	1.5329	.0131	—
	1.5127	.0175	—	Telescope (Sb) flint	1.5286	.0194	2.50
	1.5215	.0168	2.50	Borosilicate flint .	1.5503	.0203	2.81
U.V. crown 3199 .	1.5035	.0155	—		1.5753	.0218	2.90
Borosilicate crown	1.4944	.0151	2.33		1.5489	.0187	—
	1.5141	.0156	2.47		1.5825	.0216	—
Barium crown .	1.5726	.0174	3.21	Barium flint . .	1.5848	.0189	—
	1.6120	.0180	—		1.6235	.0256	3.67
Heavy barium crown	1.6130	.0178	3.60		1.6570	.0276	3.95
Flints—					1.7174	.0340	4.49
(Silicate) flint .	1.5794	.0244	3.25	Heavy flint . .	1.7782	.0378	4.99
	1.6138	.0271	3.58		1.9044	.0461	5.92
	1.6489	.0296	3.87		1.9625	.0508	—

SPECTROSCOPY

It is now agreed that the use of the diffraction-grating in fundamental work must be limited to interpolation between standard wave-lengths obtained by other means. The accepted standard lines are three in the spectrum of cadmium. Their wave-lengths (λ) obtained by interference methods, and measured (by direct comparison with the standard metre at Paris) in dry air at 15° C. (H-scale) and 760 mms. mercury pressure, are given below in tenth-metres ($= 10^{-8}$ cm. = 1 Angström unit). (See Michelson's "Light Waves and their Uses.") [$\mu = 10^{-4}$ cm.; $\mu\mu = 10^{-7}$ cm.]

Observer.	λ Cd red.	λ Cd green.	λ Cd blue.
Michelson and Benoit, 1894	6438.4700	5085.8218	4799.9085
Benoit, Fabry, and Perot, 1907 . . .	6438.4702	—	—

The following values (all in tenth-metres) are of course only approximate:—

Hertzian Waves.	Infra-red.	Red.	Orange.	Yellow.	Green.	Blue.	Violet.	Ultra-violet.	
$10^{14} - 4 \times 10^7$	3.1×10^8	§ 7700	6470	5880	5500	4920	4550	3600	600

STANDARD LINES—IRON ARC SPECTRUM

Obtained by an interference method, and based on Benoit, Fabry, and Perot's value for the wave-length of the red line of cadmium. The wave-lengths below are given in tenth-metres (10^{-8} cm.), measured in dry air at 15° (H-scale) and 760 mms. mercury. (Buisson and Fabry, *Compt. Rend.*, 1907 and 1909.)

2373.737	2987.293	3724.379	4352.741	4878.226	5405.780	5952.739
2413.310	3030.152	3753.615	4375.935	4903.324	5434.530	6003.039
2435.159 *	3075.725	3805.346	4427.314	4919.006	5455.616	6027.059
2506.904 *	3125.661	3843.261	4466.554	4966.104	5497.521	6065.493
2528.516 *	3175.447	3865.526	4494.572	5001.880	5506.783	6137.700
2562.541	3225.790	3906.481	4531.155	5012.072	5535.418	6191.569
2588.016	3271.003	3935.818	4547.854	5049.827	5569.632	6230.732
2628.296	3323.739	3977.745	4592.658	5083.343	5586.770	6265.147
2679.065	3370.789	4021.872	4602.944	5110.415	5615.658	6318.029
2714.419	3399.337	4076.641	4647.437	5127.364	5658.835	6335.343
2739.550	3445.155	4118.552	4678.855	5167.492	5709.396	6393.612
2778.225	3485.344	4134.685	4707.287	5192.362	5760.843 †	6430.859
2813.290	3513.820	4147.677	4736.785	5232.958	5763.013	6494.994
2851.800	3556.879	4191.441	4754.046 †	5266.568	5805.211 †	† Si.
2874.176	3606.681	4233.615	4789.657	5302.316	5857.760 †	† Mn.
2912.157	3640.391	4282.407	4823.521 †	5324.196	5892.882 †	† Ni.
2941.347	3677.628	4315.089	4859.756	5371.498	5934.683	—

CHIEF ABSORPTION (FRAUNHOFER) LINES IN SOLAR SPECTRUM

Rowland's wave-lengths corrected approximately by the use of Fabry and Perot's results, measured in tenth-metres (10^{-8} cm.) in air at 20° and 760 mms. Owing to atmospheric absorption, the sun's spectrum extends only to about wave-length 3000.

Line.	Subst.	Rel. Intens.	Line.	Subst.	Rel. Intens.	Line.	Subst.	Rel. Intens.
3047.5	Fe	20	L 3820.4	Fe-C	25	(H γ) 4340.4	H	20
3057.3	Ti-Fe	20	3825.8	Fe	20	F 4861.37	H (a)	30
3059.0	Fe	20	3838.2	Mg-C	25	b $_2$ 5172.7	Mg	20
O 3440.6	Fe	20	3859.8	Fe-C	20	b $_1$ 5178.22	Mg	30
3441.0	Fe	15	K 3933.6	Ca	1000	E 5269.56	Fe	8
3524.5	Ni	20	3961.5	Al	20	(D $_2$) 5875.62 †	He	—
N 3581.2	Fe	30	H 3968.4	Ca	700	D $_2$ 5889.97	Na	30
3608.8	Fe	20	4045.8	Fe	30	D $_1$ 5895.93	Na	20
3618.7	Fe	20	4063.6	Fe	20	C 6562.8	H (a)	40
M 3719.9	Fe	40	(H δ) 4101.8	H	40	B 6867.3	†	6
3734.8	Fe	40	4226.7	Ca	20	A 7661 *	†	—
3737.1	Fe	30	G 4307.9	Fe	6	Z 8228 *	—	—

* Langley, 1900.

† Oxygen in earth's atmos.

† Emission line in chromosphere alone.

§ Wood, 1911.

|| X and γ rays 8.4 to 0.07.

EMISSION SPECTRA

EMISSION SPECTRA OF SOLIDS

For a fuller treatment of wave-lengths see Watts' "Index of Spectra" and appendices, Kayser's "Handbuch der Spectroscopie," Hagenbach and Konen's "Atlas of Emission Spectra," 1905. For recent work consult the *Astrophysical Journal*. The wave-lengths below are measured in tenth-metres (10^{-8} cm.) in air at 15° C. and 760 mms. The visible spectrum colours are indicated—*r*, *o*, *y*, *g*, *b*, *v*.

The brightest lines are emphasized and the approximate boundary of the ultra-violet region is indicated thus

ALUMINIUM (arc).	CADMIUM (contd.)	CALCIUM (contd.)	MAGNESIUM (contd.)	RADIUM (contd.)	SODIUM (NaCl in flame).
3083	4413 <i>b</i>	6122 <i>o</i>	3832	4683 <i>v</i>	Fabry and
3093	4678 <i>b</i>	6162 <i>o</i>	3838	4826 <i>b</i>	Perot, 1902 ;
.....	4799-908 <i>b</i>	6140 <i>o</i>	5168 <i>g</i>	5210 <i>g</i>	Rayleigh, '06.
3944 <i>v</i>	5085-822 <i>g</i>	6163 <i>o</i>	(<i>b</i> ₂) 5173 <i>g</i>	5360 <i>g</i>	(D ₂) 5889-9650
3962 <i>v</i>	5338 <i>g</i>	6500 <i>r</i>	5184 <i>g</i>	5655 <i>y</i>	(D ₁) 5895-9320
4663 <i>b</i>	5379 <i>g</i>		5529 <i>y</i>	5685 <i>y</i>	
5057 <i>g</i>	6438-470 <i>r</i>	COPPER (are in vacuo).		6210 <i>o</i> ³	STRONTIUM (SrCl ₂ in flame).
5696 <i>y</i>		Fabry and	MERCURY (Mercury lamp).	6216 <i>o</i> ³	(SrCl ₂ in flame).
5723 <i>y</i>	CAESIUM (CsCl in flame).	Perot, 1902.	Stiles, <i>Astro.</i>	6228 <i>o</i> ³	Band spectr'm
5	3611-8	3248	<i>Journ.</i> , 1909.	6247 <i>o</i> ³	with lines at
Full of bands, some diffuse, and some resolvable.	3617	3274	3126	6250 <i>o</i> ³	4607-5 <i>b</i>
	3877	3131	6260 <i>o</i> ³	6387 <i>o</i>
	3889	4023 <i>v</i>	3650	6285 <i>o</i> ³	THALLIUM (Tl or TlCl ₂ in flame).
3501	4555 <i>b</i>	4063 <i>v</i>	4046-8 <i>v</i>	6329 <i>o</i> ³	5350-7 <i>g</i>
.....	4593 <i>b</i>	5105-543 <i>g</i>	4078-1 <i>v</i>	6349 <i>o</i>	
3910 <i>v</i>	5664 <i>y</i>	5153-251 <i>g</i>	4358-343 <i>v</i> ²	{ 6530 <i>r</i> ³	
3994 <i>v</i>	5845 <i>y</i>	5218-202 <i>g</i>	4916-4 <i>b</i> <i>g</i>	to	
4131 <i>v</i>	6011 <i>o</i>	5700 <i>y</i>	4959-7 <i>g</i>	{ 6700 <i>r</i> ³	
4131 <i>v</i>	6213 <i>o</i>	5782-090 <i>y</i>	5460-742 <i>g</i> ²	6653 <i>r</i>	
4554 <i>b</i>	6724 <i>r</i>	5782-159 <i>y</i>	5769-598 <i>y</i> ²	³ Bands.	TIN (spark).
4934 <i>g</i>	6974 <i>r</i>	INDIUM (In(OH) ₂ in flame).	5790-659 <i>y</i> ²	RUBIDIUM (RbCl in flame).	3009
m 5536 <i>gy</i>		6152 <i>o</i>	6152 <i>o</i>	3349	3034
5778 <i>y</i>	CALCIUM (CaCl ₂ in flame).	6232-0 <i>o</i>	6232-0 <i>o</i>	3351	3175
5854 <i>y</i>	Bands pre- dominate ; line at	4102 <i>v</i>	² Fabry and Perot, 1902, and Rayleigh, 1906.	3587	3262
6142 <i>o</i>	4227	4511 <i>v</i>	POTASSIUM (KCl in flame).	3592	3283
6497 <i>r</i>	(Flame arc).	IRON (see p. 75).	3446	3331
BORON (Boric acid in flame).	3362	LITHIUM (LiCl in flame).	3447	4202 <i>v</i>	3596
Diffuse maxima at	3644	4132 <i>v</i>	4216 <i>v</i>	3746
4500 <i>b</i>	(K) 3934 <i>v</i>	4602 <i>b</i>	4044 <i>v</i>	5618 <i>y</i>	4525 <i>v</i>
4700 <i>b</i>	(H) 3968 <i>v</i>	6104 <i>o</i>	4047 <i>v</i>	5724 <i>y</i>	5563 <i>y</i>
4900 <i>b</i>	4227 <i>v</i>	6707-846 <i>r</i> ¹	5802 <i>y</i>	6207 <i>o</i>	5589 <i>y</i>
5200 <i>g</i>	4303 <i>b</i>	¹ Fabry and Perot, 1902.	7668 <i>r</i>	6298-7	5799 <i>y</i>
5450 <i>g</i>	4426 <i>b</i>	MAGNESIUM (arc).	7702 <i>r</i>		6453 <i>o</i>
5800 <i>y</i>	4435 <i>b</i>	3091		SILVER (are in vacuo).	ZINC (are in vacuo).
6000 <i>o</i>	4455 <i>b</i>	3093	RADIUM (RaBr ₂ in flame).	4055 <i>v</i>	3036
CADMIUM (arc).	4586 <i>b</i>	3097	Runge and Precht, 1903.	4212 <i>v</i>	3072
3261	4878 <i>b</i>	3330	3650	4669 <i>b</i>	3345
3404	5270 <i>g</i>	3332	3815	5209-081 <i>g</i> ⁴	4680-138 <i>b</i> ⁵
3466	5350 <i>g</i>	3337	5465-489 <i>g</i> ⁴	4722-164 <i>b</i> ⁵
3611	5589 <i>y</i>	3830	4341 <i>v</i>	5472 <i>g</i>	4810-535 <i>b</i> ⁵
.....	5595 <i>y</i>			5623 <i>g</i>	4912 <i>b</i>
3982 <i>v</i>	5858 <i>y</i>			⁴ Fabry and Perot, 1902	4925 <i>g</i> ⁶
					6103 <i>o</i>
					6362-345 <i>o</i> ⁵
					⁵ Fabry and Perot, 1902.

EMISSION AND ABSORPTION SPECTRA

EMISSION SPECTRA OF GASES

The gases are all in vacuum tubes (2-4 mms. press.); only the brightest lines are given. The visible spectrum colours are indicated—*r*, *o*, *y*, *g*, *b*, *v*.

See the general remarks on last page.

ARGON, Red spectrum (small current density).	CARBON MONOXIDE or DIOXIDE (of common oc- currence in many vacuum- tube spectra). Numerous bands shaded towards violet edges at	HYDROGEN Elementary spec- trum.	NEON (<i>contd.</i>)	NITROGEN (<i>contd.</i>)
4159 <i>v</i>		3750	5853 <i>y</i>	5804 <i>y</i>
4192 <i>v</i>		3771	5882 <i>o</i>	5854 <i>y</i>
4198 <i>v</i>		3798	5945 <i>o</i>	5906 <i>o</i>
4201 <i>v</i>		3836	5976 <i>o</i>	5959 <i>o</i>
4259 <i>b</i>		3889	6030 <i>o</i>	6013 <i>o</i>
4300 <i>b</i>		6075 <i>o</i>	6069 <i>o</i>
4334 <i>b</i>		3970 <i>v</i>	6096 <i>o</i>	With large cur- rent densities, N gives a line spectrum.
4511 <i>b</i>	3590 (CN)	4102 (8) <i>v</i>	6129 <i>o</i>	
4703 <i>b</i>	3884 (CN)	4340 (γ) <i>b</i>	6143 <i>o</i>	
5452 <i>g</i>	(F) 4861 (8) <i>gb</i>	6164 <i>o</i>	OXYGEN Elementary line spectrum.
5607 <i>y</i>	4123 <i>v</i>	(C) 6563 (α) <i>r</i>	6182 <i>o</i>	
5912 <i>o</i>	4216 (CN) <i>v</i>	For very short wave-lengths	6217 <i>o</i>	
6031 <i>o</i>	4393 <i>b</i>	(1030-1675) see	6267 <i>o</i>	3919 3973 4070 <i>v</i>
6059 <i>o</i>	4511 <i>b</i>	Lyman, <i>Astro.</i>	6305 <i>o</i>	
	4735 (C) <i>b</i>	<i>Journ.</i> , 1906.	6383 <i>o</i>	
	4835	Secondary spec- trum	6402 <i>o</i>	Diffuse maxima at
	5165 (C) <i>g</i>	(see Watson, <i>Proc. Roy. Soc.</i> , 1909).	6507 <i>r</i>	
	5198 <i>g</i>			
	5610 <i>y</i>			There are three other oxygen spectra: con- tinuous, band, and series spectra.
	6079 <i>o</i>			
Blue spectrum (large current density).	HELIUM Rayleigh, 1908.	KRYPTON AND XENON <i>Brit. Ass. Rep.</i> , 1905.	NITROGEN Band spectrum from positive column. Many bands all made up of fine lines. From 3000 to 4574 the edges occur at inter- vals of about 60 Å.U. Other bands have edges at	5335 <i>g</i>
3583	3188	NEON Baly, <i>Phil.</i>	4648 <i>b</i>	5440 <i>g</i>
4072 <i>v</i>	<i>Trans.</i> , 1903.	4666 <i>b</i>	6110 <i>o</i>
4104 <i>v</i>	3889 <i>v</i>	Very rich in	4723 <i>b</i>	6170 <i>o</i>
4228 <i>v</i>	4026 <i>v</i>	red rays.	4813 <i>b</i>	RADIUM EMANA- TION Royds, <i>Phil.</i> <i>Mag.</i> , 1909.
4331 <i>b</i>	4471·482 <i>b</i>		5340 <i>g</i>	
4348 <i>b</i>	4713·144 <i>b</i>		5614 <i>y</i>	
4426 <i>b</i>	4921·930 <i>gb</i>		5755 <i>y</i>	
4430 <i>b</i>	5015·680 <i>g</i>			
4431 <i>b</i>	(D ₃) 5875·625 <i>y</i>			
4610 <i>b</i>	6678·150 <i>r</i>			
4806 <i>b</i>	7065·200 <i>r</i>			

ABSORPTION SPECTRA

For wave-lengths of the Fraunhofer lines in the sun's spectrum, see p. 75.

Among the enormous literature on absorption spectra, reference may be made to Kayser's "Handbuch der Spectroscopie," Baly's "Spectroscopy," Vogel's "Praktische Spectralanalyse," the writings of Prof. Hartley, Jones and Anderson's "Absorption Spectra of Solutions," 1909, Smiles' "Chemical Constitution and Physical Properties," and the British Association Reports of 1901 *et seq.*

Convenient substances which show good absorption spectra are—neodymium and praseodymium salts and didymium glass (which yield some extremely narrow absorption lines), iodine vapour, nitrogen peroxide, chlorine, chlorophyll, blood, and potassium permanganate solution.

OPTICAL ROTATIONS OF PURE LIQUIDS AND SOLUTIONS

A_t = the rotation in degrees (for light of some given wave-length) of the plane of polarization by a liquid when at the temperature t° C.

l_t = the length of the column of liquid in **decimetres** (*i.e.* 10 cms.).

ρ = the number of grams of active substance in 100 **grams of solution**.

q = $(100 - \rho)$ = the percentage (by weight) of inactive solvent in the solution.

ρ_t = the density in grams per c.c. of the liquid or solution at t° .

$c_t = \rho \rho_t$ = the concentration expressed as grams of active substance per 100 **c.cs. of solution** at t° .

$[\alpha]_t$ = the **specific rotation** (at t°) = $\frac{\text{rotation per decimetre of sol.}}{\text{grams of active substance per c.c. of sol.}}$

For a pure liquid $[\alpha]_t = \frac{A_t}{l_t \rho_t}$.

For an active substance in solution $[\alpha]_t = \frac{A_t}{l_t} \left(\frac{\rho}{\rho + q} \rho_t \right) = \frac{100 A_t}{l_t \rho \rho_t} = \frac{100 A_t}{l_t c_t}$, since $(\rho + q) = 100$.

The rotation depends on the wave-length of the light used; it increases as the wave-length (λ) diminishes ($\alpha \propto \frac{1}{\lambda^2}$ approx.). α also varies with the nature of the inactive solvent and with the concentration of the solution.

The rotation is called positive or right-handed (*dextro*, d) if the plane of polarization appears to be rotated in an anti-clockwise direction when looking through the liquid **away** from the source of light. The contrary rotation is called *laevo* (l). The **molecular rotation** is the specific rotation multiplied by the molecular weight.

$[\alpha]_{20}^D$ indicates that the specific rotation is measured at 20° C. using sodium (D) light.

(See Landolt's "Optical Rotations of Organic Substances and their Practical Application.")

Optically Active Substance.	Solvent.	Conditions.	Specific Rotation $[\alpha]_t$
Cane Sugar or Candy (d), $C_{12}H_{22}O_{11}$ (Landolt, 1888; Pellat, 1901)	water	$c = 4$ to 28	$[\alpha]_{20}^D = +66.67 - .0095c$
		$t = 14^\circ$ to 30° C.	$[\alpha]_t^D = [\alpha]_{20}^D \{1 - .00037(t - 20)\}$
Invert Sugar (l),* $C_6H_{12}O_6$ \equiv 1 mol. of dextrose + 1 mol. of levulose (Gubbe, 1885)	water	$c = 9$ to 35	$[\alpha]_{20}^D = -19.7 - .036c$
		$t = 3^\circ$ to 30° C.	$[\alpha]_t^D = [\alpha]_{20}^D + .304(t - 20) + .00165(t - 20)^2$
Dextrose (d - glucose), $C_6H_{12}O_6$ (Parcus and Tollens, 1890; Tollens, 1884)	water	$c = 9.1$	$[\alpha]_{20}^D = +105.02$ after 5.5 mins. (α modification) $= +52.5$ after 6 hrs. (β modification)
	water	$\rho = 1$ to 18	$[\alpha]_{20}^D = +52.5 + .025\rho$
l - Glucose , $C_6H_{12}O_6$ (Fischer, 1890)	water	$\rho = 4$	$[\alpha]_{20}^D = -94.4$ after 7 mins. $= -51.4$ after 7 hrs.
Levulose (l) (fruit sugar), $C_6H_{12}O_6$ (Parcus and Tollens, 1890; Ost, 1891)	water	$c = 10$	$[\alpha]_{20}^D \doteq -104^\circ$ after 6 mins. $= -92^\circ$ after 33 mins.
	water	$\rho = 2$ to 31	$[\alpha]_{20}^D = -91.9 - .11\rho$

* The molecular weight of cane-sugar is 342; which, after conversion to invert sugar, becomes 360. Hence the new concentration of the invert sugar solution is $\frac{342}{360}c$, where c is the number of grams of cane-sugar in 100 c.cs. of the original solution.

Optically Active Substance.	Solvent.	Conditions.	Specific Rotation $[\alpha]_t$
Galactose (d), $C_6H_{12}O_6$ (Meissl, 1880)	water	$\phi = 4$ to 36 $t = 10^\circ$ to $30^\circ C.$	$[\alpha]_t^D = +83^\circ 9 + \cdot 078\phi$ $- \cdot 21t$
Ord. Tartaric acid (d), $H_2C_4H_4O_6$	water	—	$[\alpha]_{20}^D = +15^\circ 06 - \cdot 131c$
Potassium tartrate (d), $K_2C_4H_4O_6$ (Thomsen, 1886)	water	$c = 8$ to 50	$[\alpha]_{20}^D = +27^\circ 14 + \cdot 0792c$ $- \cdot 00094c^2$
Rochelle salt (d), $KNaC_4H_4O_6$	water	—	$[\alpha]_{20}^D = +29^\circ 73 - \cdot 0078c$
<i>l</i> - Turpentine , $C_{10}H_{16}$ (Gernez, 1864; Landolt, 1877)	pure liquid	—	$[\alpha]_{20}^D = -37^\circ$
	vapour	at 761·7 mms.	$[\alpha]_{168}^D = -35^\circ 5$ for mean yellow
	alcohol ($\rho_{20} = \cdot 796$)	$q = 0$ to 90	$[\alpha]_{20}^D = -37^\circ - \cdot 00482q$ $- \cdot 00013q^2$
	benzene	$q = 0$ to 91	$[\alpha]_{20}^D = -37^\circ - \cdot 0265q$
	paraffin oil	Within wide limits $[\alpha]$ increases with the percentage of paraffin.	
Quinine sulphate (l), $C_{20}H_{24}N_2O_2 \cdot H_2SO_4$ (Oudemans, 1876)	water	c about 1·6 % of alkaloid (calculated)	Salt $[\alpha]_{17}^D = -214^\circ$ Alkaloid $[\alpha]_{17}^D = -278^\circ$
Nicotine (l), $C_{10}H_{14}N_2$ (Landolt, 1877; Hein, 1898)	pure	$t = 10^\circ$ to $30^\circ C.$	$[\alpha]_{20}^D = -162^\circ$
	benzene	$\phi = 8$ to 100	$[\alpha]_{20}^D = -164^\circ$
	water	$\phi = 1$ to 16	$[\alpha]_{20}^D = -77^\circ$
Ethyl malate (l), $(C_2H_5)_2C_4H_4O_5$ (Purdie & Williamson, '96)	pure liquid	—	$[\alpha]_{11}^D = -10^\circ 3$ to $-12^\circ 4$
Camphor (d), $C_{10}H_{16}O$ (Landolt, 1877; Rim-bach, 1892)	alcohol	$q = 45$ to 91	$[\alpha]_{20}^D = +54^\circ 4 - \cdot 135q$
	benzene	$q = 47$ to 90	$[\alpha]_{20}^D = +56^\circ - \cdot 166q$

OPTICAL ROTATION AND WAVE-LENGTH

Wave-length (λ) in 10^{-8} cm.	Specific Rotation at $20^\circ C.$ $[\alpha]_{20}^\lambda$				QUARTZ AT $20^\circ C.$	
	Cane-sugar or Candy in H_2O .	Turpentine (pure liq.).	Tartaric acid in H_2O ($p = 41\%$).	Nicotine (pure liq.).	Wave-length (λ) in 10^{-8} cm.	Rotation for 1 mm. thickness.
H (C) 6563 (r)	52°·9	-29°·5	7°·75	-126°	Li 6708 (r)	16°·4
Na (D) 5893 (ϕ)	66·5	-37	8·86	-162	H (C) 6563 (r)	17·3
					Na (D) 5893 (ϕ)	21·72*
Tl 5351 (g)	81·8	-45	9·65	-207·5	Tl 5351 (g)	26·53
					H (F) 4861 (g)	32·7
H (F) 4861 (g)	100·3	-54·5	9·37	-253·5	H (δ) 4102 (b)	47·48

* For quartz at temperature t° , rotation = $21^\circ 72 \{1 + \cdot 000147(t - 20)\}$ for D line.

FARADAY EFFECT

MAGNETIC ROTATION OF POLARIZED LIGHT

This effect was discovered by Faraday in 1845. The rotation per cm. per unit magnetic field—**Verdet's constant**, $r = \alpha/(Hl)$, where α is the rotation in minutes for the substance in a magnetic field of H gauss, and l is the length of light-path parallel to the lines of force. r varies with the temperature and is roughly inversely proportional to the square of the wave-length of the light used. Films of Fe, Ni, and Co are exceptions to this rule.

If the light is travelling with the lines of force (*i.e.* from N. to S.), then the direction of rotation is positive, if the plane of polarization is rotated clockwise, to an observer looking in the direction in which the light is moving. If the light is reflected back on its path, the rotation is increased.

The **Molecular rotation** $r_m = rM/d$, where M is the molecular weight of the substance, and d is its density. r_m is an additive property in organic compounds (Perkin, *Journ. Chem. Soc.*, 1884).

The rotations below are for the sodium D line ($\lambda = 5893 \times 10^{-8}$ cm.).

(For Voigt's theory of magneto-rotation, see Schusters, "Optics," 1909. See also Becquerel's papers in *Compt. Rend.*, etc.)

Substance.	Temp.	Rotation r in mins. of arc.	Substance.	Temp.	Rotation relative to Water.
Water	0°C.	+01311, R.W.	Ethyl alcohol . . .	16.8	*8637, P.
"	20	+01312, R.W.	n. propyl alcohol . .	15.6	*9139, P.
Carbon bisulphide . .	0	+04347, R.W.	Amyl(iso) alcohol . .	19.9	*9888, P.
"	18	+04200, Ra.	Ethyl bromide . . .	19.7	1395, P.
Quartz, \perp axis . . .	20	+01368, * Bo.	" chloride	5.0	1035, P.
"	20	+01664, Bo.	" iodide	18.1	2251, P.
"	20	+1587, † Bo.	Formic acid	20.8	7990, P.
Jena (phosphate crown	18	+0161, D.B.	Acetic "	21.0	7976, P.
glass) heaviest flint .	18	+0888, D.B.	Propionic acid . . .	20.3	*8369, P.
FeCl ₃ dens. = 1.693 .	15	-2026, B.	Benzene	15	2062, B.
"	1.023	15			

* $\lambda = 6439$. † $\lambda = 2194$. B., Becquerel; Bo., Borel, 1903; D.B., Du Bois, 1894; P., Perkin; Ra., Rayleigh, 1884; R.W., Rodger and Watson, 1896.

METALLIC REFLECTION OF LIGHT

(The percentage of normally incident light reflected from different surfaces.)

The column of figures (below) in the case of **speculum metal** (7 Cu, 3 Sn) reads 30% (for $\lambda = 2510$); 51%, 56%, 64%, 67%, 71%, 89%, 94% (for $\lambda = 140,000$).

Wave-length λ in A.U. (10^{-8} cm.).	Cu.	Au.	Ni.	Pt.	Ag.	Steel.	Magna- lium.*	Glass mirror.	
								Ag back.	Hg back.
Ultra-violet {	2,510	26%	39%	38%	34%	34%	67%	—	—
	3,570	27	28	49	43	74	81	—	—
Visible {	4,200	33	29	57	52	87	83	86%†	73%†
	5,500	48	74	63	61	93	55	88	71
	7,000	83	92	69	69	95	58	83	73
Infra-red {	10,000	90	95	72	73	97	63	84	
	40,000	97	97	91	91	98	88	89	
	140,000	98	98	97	96	99	96	92	

* 69 Al, 31 Mg.
† $\lambda = 4500$.

DIOPTER

In applied optics the "power" of a lens or mirror is expressed in diopters. The number of diopters equals the reciprocal of the focal length expressed in metres.

ELECTRICAL RESISTIVITIES

Electrical specific resistances or resistivities in ohm-cms. **Conductivities** (in reciprocal ohms) are the reciprocals of resistivities. For a table of reciprocals, see p. 136.

METALS AND ALLOYS

The resistivity depends to some extent on the state of the metal. In general, cold drawing increases, while annealing diminishes the resistance. The winding of a wire into a coil increases its resistance.

For pure metals, the resistance is roughly proportional to the absolute temperature, and would apparently vanish not far from the absolute zero. This rule does not hold even approximately for alloys.

For wire resistances, see p. 83; for temperature coefficients, next page. The thermal conductivities of the same samples of many of the substances below will be found on p. 51.

Substance.	Temp.	Sp. Re.	Observer.	Substance.	Temp.	Sp. Re.	Observer.
Metals—	° C.	$\times 10^{-6}$		Metals (contd.)	° C.	$\times 10^{-6}$	
Aluminium* . . .	-160	0.81	Lees,	Platinum . . .	-203	2.4	D.&F., '96
" . . .	18	2.94	P. T., '08	" . . .	18	11.0	J. & D.,
" . . .	18	3.21	J. & D.,	" . . .	100	14.0	1900
" . . .	100	4.13	1900	Potassium . . .	0	6.64	B., '04
Antimony . . .	15	40.5	Berget, '90	Rhodium . . .	18	6.0	—
Bismuth . . .	18	119.0	J. & D.,	Silver, 99.9% . .	-160	0.56	Lees,
" . . .	100	160.3	1900	" . . .	18	1.66	1908
Cadmium, drawn .	-160	2.72	Lees, '08	" . . .	18	1.63	J. & D.,
" . . .	18	7.54	J. & D.,	" . . .	100	2.13	1900
" . . .	100	9.82	1900	Sodium . . .	0	4.74	B., 1904
Copper, drawn . .	-160	6.49	Lees, '08	Strontium . . .	20	25	M., 1857
" . . .	18	1.78	J. & D.,	Tantalum . . .	18	14.6	—
" . . .	100	2.36	1900	Tellurium . . .	20	21	M., 1858
" annealed . . .	18	1.59	Mean	Thallium, pure . .	0	17.6	D.&F., '96
Calcium . . .	20	10.5	M.&C., '05	Thorium . . .	15	40.1	Bo., '09
Cobalt . . .	20	9.71	R., 1901	Tin, drawn . . .	-160	3.5	Lees, '08
Gold . . .	-183	0.68	D.&F., '96	" . . .	18	11.3	J. & D.,
" . . .	18	2.42	J. & D.,	" . . .	100	15.3	1900
" . . .	100	3.11	1900	Tungsten . . .	25	5.0	Fink, '10
Iridium . . .	18	5.3	—	Zinc, pure . . .	-160	2.2	Lees, '08
Iron . . .	18	9.15	Mean	" . . .	18	6.1	J. & D.,
" (1%) . . .	18	12.0	J. & D.,	" . . .	100	7.9	1900
" (C.) . . .	100	16.8	1900				
" wrought . . .	-160	5.4	Lees, '08	Alloys—			
" . . .	18	13.9	J. & D.,	Brass . . .	-160	4.1	Lees,
" . . .	100	18.8	1900	" . . .	17	6.6	1908
" steel (1%) . .	18	19.9	J. & D.,	" . . .	18	6.9	Mean
" (C.) . . .	100	25.6	1900	Constantan . . .	18	49.0	J. & D.,
Lead, drawn . . .	-160	7.43	Lees, '08	(Eureka)§ . . .	100	49.1	1900
" . . .	18	20.8	J. & D.,	German silver .	18	16.40	Mean
" . . .	100	27.7	1900	" . . .	0	26.6	Lorenz,
Lithium . . .	0	8.4	B., '04	" . . .	100	27.6	1881
Magnesium . . .	0	4.35	D. & F.	Manganin¶ . . .	-160	43.13	Lees,
Mercury . . .	0	94.07	See	" . . .	18	44.50	1908
" . . .	20	95.76	pp. 6, 82.	" . . .	18	42.05	J. & D.,
Molybdenum . . .	25	4.1	Fink, '10	" . . .	100	42.11	1900
Nickel . . .	-160	5.9	Lees, '08	Phosphor-bronze .	18	5.10	Mean
" (97%) . . .	18	11.8	J. & D.,	Platinoid . . .	-160	32.5	Lees,
" (Ni) . . .	100	15.7	1900	" . . .	18	34.4	1908
Osmium . . .	20	9.5	Blair, '05	90 Pt, 10 Rh . . .	0	21.1	D.&F., '96
Palladium . . .	18	10.7	J. & D.,	67 Pt, 33 Ag . . .	0	24.2	—
" . . .	100	13.8	1900				

* 99% Al. † 1% C, 2% Si, 1% Mn.

§ 60 Cu, 40 Ni. || 62 Cu, 15 Ni, 22 Zn.

† 70 Cu, 30 Zn.

¶ 84 Cu, 4 Ni, 12 Mn.

B., Bernini; Bo., Bolton; D. & F., Dewar & Fleming; J. & D., Jaeger and Diesselhorst; M., Matthiessen; M. & C., Moissan & Chavanne; R., Reichardt; P. T., Phil. Trans.

RESISTIVITIES

ELECTRICAL RESISTIVITIES (*contd.*)

NON-METALS AND INSULATORS

The resistivities are in ohm-cms. at room temperatures unless otherwise stated. The values for insulators naturally vary widely, and the figures below are merely typical and are probably, in many cases, nothing more than the resistances of the surfaces. For a discussion of some electrical insulators, see Kaye, *Proc. Phy. Soc. Lond.*, 1911.

Substance.	Sp. Re.	Substance.	Sp. Re.	Substance.	Sp. Re.
Gas carbon . . .	{ '004 to '007	Sulphur, 70° . .	4. 10 ¹⁵	Guttapercha . .	2. 10 ⁹
Graphite . . .	'003	Ebonite . . .	2. 10 ¹⁵	Mica	9. 10 ¹⁶
C. lamp filament	'004	Glass, soda-lime *	5. 10 ¹¹	Paraffin wax . .	3. 10 ¹⁸
Selenium † (1907)	2. 10 ¹⁶	" Jena, com- bustion *	> 2. 10 ¹⁴	Porcelain, 50° . .	2. 10 ¹³
Silicon §	'06	" conducting†	5. 10 ⁸	Quartz	1'2. 10 ¹⁴
				Fused silica * . .	> 2. 10 ¹⁴

* National Physical Laboratory.

† Phillips.

‡ In dark.

§ Wick, 1908.

TEMPERATURE COEFFICIENTS OF RESISTANCE

To represent accurately over any considerable range the variation of electrical resistance (R) with temperature (t) requires for almost all substances a parabolic or cubic equation in t . But if the temperature interval is not large, a linear equation $R_t = R_0(1 + \alpha t)$ may be employed; and this gives a definition of the mean temperature coefficient (α) over that temperature range. The table of resistivities above will readily yield the associated values of α . The coefficients given below are average ones.

Substance.	Temp.	α	Substance.	Temp.	α
Metals—		$\times 10^{-4}$	Metals (<i>contd.</i>)—		$\times 10^{-4}$
Aluminium	18-100	38	Silver	0-100	40
Bismuth	18	42	Tantalum	0-100	33
Cadmium	18-100	40	Tin	0-100	45
Copper *	18	42'8	Tungsten (1910) . .	0-170	51
Cobalt	0-160	33	Zinc	18-100	37
Gold	0-100	40			
Iron, pure	18	62	Alloys—		
Steel	18	16-42	Brass	18	10 ‡
Lead	18	43	Constantan (Eureka) .	18	{ - '4 to + '1 ‡
Mercury †	0-24	9'0	German silver	18	2'3-6
Nickel, electrolytic .	0-100	62	Manganin §	20	'02-5 ‡
" commercial	0-1000	27	Platinoid	18	2'5
Palladium	18-100	37	90 Pt, 10 Ir	16	15
Platinum	-100-0	35	90 Pt, 10 Rh	15	17
"	0-100	38	Platinum-silver (coils)	16	2'4-3'3
Molybdenum (1910) .	0-170	50			

* High conductivity annealed (N. P. L.), 1904.
from 30° C. to 40° C.

‡ N. P. L.

† $R_t = R_0(1 + '0,88t + '0,1t^2)$ —Smith
§ Most samples of manganin have a zero temp. coeff. at

STANDARD WIRE GAUGE

The sizes of wires are ordinarily expressed by an arbitrary series of numbers. There are, unfortunately, four or five independent systems of numbering, so that the wire gauge used must be specified. The following are English Legal Standard wire gauge values. (See Foster's "Electrical Engineers' Pocket Book.")

Size.			Diameter.			Size.			Diameter.		
S.W.G.	Mm.	Inch.	S.W.G.	Mm.	Inch.	S.W.G.	Mm.	Inch.	S.W.G.	Mm.	Inch.
6	4.88	.192	20	.914	.036	34	.234	.0092			
8	4.06	.160	22	.711	.028	36	.193	.0076			
10	3.25	.128	24	.559	.022	38	.152	.0060			
12	2.64	.104	26	.457	.018	40	.122	.0048			
14	2.03	.080	28	.376	.0148	42	.102	.0040			
16	1.63	.064	30	.315	.0124	44	.081	.0032			
18	1.22	.048	32	.274	.0108	46	.061	.0024			

WIRE RESISTANCES

Average values in ohms per metre at 15° C. The **safe currents** for copper (high conductivity annealed commercial) are calculated at the rate of about 270 amps./cm.² for No. 12 wire, 430 amps./cm.² for No. 22 wire, and 500 amps./cm.² for smaller diameters. Larger current densities than these are allowed in the revised "Wiring Rules" of the Institution of Electrical Engineers. Eureka is practically identical with constantan.

The average **temperature coefficient** of resistance of copper is .00428; of nickel, .0027; of manganin, .0001; of German silver, .00044; of Eureka, -.00002; of platinoid, .00025 per degree Centigrade. The values for the alloys may vary considerably. The **composition** of manganin is 84Cu, 4Ni, 12Mn; of German silver, 60Cu, 15Ni, 25Zn; of Eureka, c. 60Cu, 40Ni. Platinoid is said to be German silver with a little tungsten. For specific resistances, see p. 81.

S.W.G.	COPPER.		MANGA NIN.	GERMAN SILVER.	S.W.G.	COPPER.		MANGA NIN.	GERMAN SILVER.
	Ohms per metre.	Safe current.	Ohms per metre.	Ohms per metre.		Ohms per metre.	Safe current.	Ohms per metre.	Ohms per metre.
		amps.					amp.		
12	.0032	15.0	.077	.041	30	.222	.4	5.45	2.90
14	.0054	9.8	.131	.070	32	.293	.3	7.18	3.83
16	.0083	6.8	.204	.109	34	.404	.2	9.90	5.27
18	.0148	4.2	.361	.193	36	.590	.15	14.5	7.74
20	.0260	2.6	.645	.345	38	.950	.1	23.2	12.4
22	.0435	1.7	1.07	.57	40	1.48	.06	36.3	19.4
24	.070	1.1	1.73	.92	42	2.10	.05	53.4	27.8
26	.105	.7	2.58	1.38	44	3.30	.03	81.7	43.5
28	.155	.5	3.82	2.02	46	5.90	.02	145.5	77.4

EUREKA or CONSTANTAN.

S.W.G.	Ohms per metre.	20° C. temp.- rise caused by	S.W.G.	Ohms per metre.	20° C. temp.- rise caused by	S.W.G.	Ohms per metre.	S.W.G.	Ohms per metre.
		amps.			amps.				
12	.086	12.2	20	.722	1.5	20	.622	28	3.69
14	.146	8.2	22	1.20	.7	22	1.03	30	5.25
16	.228	4.9	24	1.93	.3	24	1.67	32	6.81
18	.405	2.7	26	2.89	.1	26	2.50	34	9.55

PLATINOID (Martino's).

S.W.G.	Ohms per metre.	20° C. temp.- rise caused by	S.W.G.	Ohms per metre.	20° C. temp.- rise caused by	S.W.G.	Ohms per metre.	S.W.G.	Ohms per metre.
		amps.			amps.				
12	.086	12.2	20	.722	1.5	20	.622	28	3.69
14	.146	8.2	22	1.20	.7	22	1.03	30	5.25
16	.228	4.9	24	1.93	.3	24	1.67	32	6.81
18	.405	2.7	26	2.89	.1	26	2.50	34	9.55

FUSES

The fusing currents are for wires mounted horizontally.

	Fusing current.	1 amp.	3	5	10	20	30	40	50
Tin	S.W.G.	37	28	24	21	18	16	14	13
Copper	S.W.G.	47	41	38	33	28	25	23	22

IONIC DISSOCIATION THEORY

On the Dissociation Theory (Arrhenius, 1887), the solute is dissociated into electrically positive cations and negative anions. For example, KCl in water exists as KCl, K^+ , Cl^- ; sulphuric acid as H_2SO_4 , H^+ , H^- , SO_4^{++} , HSO_4^+ . Probably, in many cases, these ions are attached to molecules of solvent. **The degree of dissociation** α = (number of dissociated solute molecules)/(total number of solute molecules). α is deduced from the osmotic pressure of the solution, and from its electric conductivity at different dilutions. The osmotic pressure is determined (1) directly, (2) from the raising of the boiling-point, and (3) from the depression of the freezing-point of the solvent by the presence of the solute. The equivalent conductivity (Λ) for different concentrations of any dilute solution is assumed to be proportional to the number of ions present. Λ approaches asymptotically a limiting conductivity (Λ_∞) for extreme dilutions, a state of things when, on this theory, the solute is completely dissociated. $\Lambda_m/\Lambda_\infty = \alpha$ for the equivalent concentration m . The cation and anion with their charges $+e$ and $-e$ (for monovalent ions) move in unit electric field in opposite directions with speeds or **mobilities** u_+ and u_- . The electrolytic current also obeys Ohm's Law, so that $X\kappa = (u_+ + u_-)ne$ (Kohlrausch, 1879), where X is the potential gradient in volts per cm., n the number of +ive or -ive ions per c.c., κ the conductivity of the solution in ohm $^{-1}$ cm. $^{-1}$. This becomes $u_+ + u_- = 1.037 \times 10^{-5} \Lambda$ cm./sec., since $\kappa/n = \Lambda/N$, and $Ne = 96,740$ coulombs per gm. equivalent of ions.

The mobility of electrolytic ions has been directly observed by Lodge (1886), Whetham, Orme Masson, and D. B. Steele. The ratio $u_-/(u_+ + u_-) = n$ is for the negative ion, the **migration ratio** or transport number of Hittorf (1853-9). n can be determined, when complex ions are absent, from the change of concentration at the anode and cathode during electrolysis. The **mobility** of certain organic ions is approximately inversely proportional to their linear dimension a (Laby and Carse). The existence of this relation of Ohm's Law and of a relation between the viscosity (η) of the solvent and the ionic mobilities (Kohlrausch, Hosking, and Lyle) indicates that the motion of the ion through the solution may follow Stokes' Law ($v = F/6\pi\eta a$, where F is the driving force), with the numerical constant, 6π , possibly changed.

The dissociation theory postulates the conditions existing in very dilute solutions. The rôle of the medium is rather neglected (Lowry, *Science Progress*, 1908). The dissociation should be large for a solvent with a high dielectric constant, for then the attraction between the cation and anion is small (Thomson and Nernst). This is generally true (Walden).

(Kohlrausch and Holborn, "Leitvermögen der Elektrolyten;" Whetham's "Theory of Solution")

MIGRATION RATIOS

Hittorf's migration ratio or transport number of the anion, $n = u_-/(u_+ + u_-)$; m = equivalent concentration per litre; t° = temp. of observation.

Solute.	$t^\circ C.$	Conc. m .	Ratio n .	Solute.	$t^\circ C.$	Conc. m .	Ratio n .	Solute.	$t^\circ C.$	Conc. m .	Ratio n .
KCl	—	.003	.505, S.D.	AgNO ₃	17°	.4 to .02	.526, H.	CuSO ₄	18°	{.08 to .02}	.625, M.
KBr	18°	{.03 to .01}	.504, B.	NH ₄ Cl	20	.05	.507, Be.				
KI	25°	.05	.505, Be.	TiCl ₃	22	.01	.516, Be.	HCl	10	{.05 to .02}	.159, N.S.
KNO ₃	8	.1	.497, H.	CaCl ₂	21	.005	.562, S.D.				
NaCl	18	{.03 to .009}	.604, B.	SrCl ₂	.21	.01	.56, Be.	HNO ₃	18	.25	.17
NaNO ₃	19	.05	.629, Be.	BaCl ₂	18	.01	.55	H ₂ SO ₄	11	.05	.17, Be.
LiCl	18	{.03 to .008}	.67	MgCl ₂	21	.05	.615, Be.	KOH	—	.1	.74
				ZnSO ₄	21	.05	.64, H.	NaOH	25	.04	.3, Be.
								NH ₃	21	.05	.56, Be.
				CdBr ₂	18	{.12 to .007}	.57	Ag ₂ C ₂ H ₃ O ₂	25	.01	.376, L.N.

B, Bogdan; Be, Bein; H., Hittorf; L.N., Löb and Nernst; M., Metelka; N.S., Noyes and Sammet; S.D., Steele and Denison.

CONDUCTIVITY OF SOLUTIONS

ELECTRICAL CONDUCTIVITY OF SOLUTIONS

κ_{18} = specific electric conductivity (in ohms⁻¹ cm.⁻¹) of the solution at 18° C.

ρ = mass of anhydrous solute per 100 gms. of solution.

η = the number of gm. equivalents in 1 c.c. of solution. Gm. equiv. per litre = 1000 η . To find η note that $\kappa/\Delta = \eta$.

v = volume in litres containing one gm. equivalent of solute = 1/1000 η .

Δ = equivalent conductivity = κ/η , = the conductivity in reciprocal ohms of 1 gm. equiv. in solution between electrodes 1 cm. apart. The chemical equiv. of, for example, " $\frac{1}{2}\text{CaCl}_2$ " is 111.2.

Temp. coefficient = $(d\kappa/dt)/\kappa_{18}$. (See Kohlrausch and Holborn, "Das Leitvermögen der Elektrolyten" (Teubner).) K = Kohlrausch; G = Grottrian.

CONCENTRATED SOLUTIONS

ρ %	κ_{18}	$\Delta = \frac{\kappa}{\eta}$	Temp. coef.	ρ %	κ_{18}	$\Delta = \frac{\kappa}{\eta}$	Temp. coef.	ρ %	κ_{18}	$\Delta = \frac{\kappa}{\eta}$	Temp. coef.	ρ %	κ_{18}	$\Delta = \frac{\kappa}{\eta}$	Temp. coef.
1 KCl (K.G.).				$\frac{1}{2}$ CdCl ₂ (G.) (contd.).				1 HCl (K.).				$\frac{1}{2}$ H ₂ SO ₄ (K.) (contd.).			
5	'0690	99.9	'0	30	'0282	6.5	252	5	'3948	281.0	158	70	'216	9.4	256
10	'1359	95.2	188	50	'0137	1.49	353	10	'6302	219.1	156	80	'110	3.9	349
15	'2020	91.5	179	1 AgNO ₃ (K.).				20	'7615	126.2	154	90	'107	3.22	320
20	'2677	88.9	168	5	'0256	83.4	218	30	'6620	69.8	152	100	'0157	—	031
21	'2810	87.5	166	10	'0476	74.3	217	40	'5152	39.1	—	1 KOH (K.).			
1 NaCl (K.G.).				15	'0683	67.9	215	1 HNO ₃ (K.G.).				4.2	'1464	188	'0
5	'0672	76	217	40	'1565	45.0	205	6.2	'312	307	147	8.4	'272	169	186
10	'1211	66.2	212	60	'2101	31.1	209	12.4	'542	257	142	12.6	'376	150	188
15	'1642	57.8	214	1 (NH ₄) ₂ SO ₄ (K.).				18.6	'690	211	137	16.8	'456	131	193
20	'1957	49.9	216	5	'0552	71.0	215	24.8	'768	169	137	29.4	'543	81	221
25	'2135	42.0	227	10	'1010	63.1	203	31	'782	133	139	42.0	'421	39	283
26.4	'2156	39.8	233	20	'1779	52.7	193	49.6	'634	61	157	1 NaOH (K.).			
$\frac{1}{2}$ CaCl ₂ (K.G.).				30	'2292	43.1	191	62	'496	36.4	157	2.5	'109	170	194
5	'0643	68.6	213	$\frac{1}{2}$ CuSO ₄ (K.).				$\frac{1}{2}$ H ₂ SO ₄ (K.).				5	'197	149	201
10	'1141	58.3	206	2.5	'0109	34.0	213	5	'208	198	121	10	'312	112	217
15	'1505	49.2	202	5	'0189	28.7	216	10	'391	180	128	15	'346	79	249
20	'1728	40.6	200	10	'0320	23.1	218	15	'543	161	136	20	'327	53	299
25	'1781	32.1	204	17.5	'0458	17.4	236	20	'653	140	145	30	'202	20	450
30	'1658	23.9	216	$\frac{1}{2}$ CdSO ₄ (G.).				25	'717	119	154	40	'116	8.1	65
35	'1366	16.1	236	1	'0042	42.9	210	30	'739	99	162	1 NH ₃ (K.).			
$\frac{1}{2}$ CdCl ₂ (G.).				5	'0146	29.0	206	35	'724	80	170	.1	'00025	4.25	246
1	'0055	50.1	222	25	'0430	13.8	223	40	'680	64	178	1.6	'00087	.93	238
10	'0241	20.2	217	36	'0421	8.25	255	50	'540	38	193	8	'00104	.23	262
								60	'373	20.3	213	30.5	'00019	'012	—

STANDARD SOLUTIONS FOR CALIBRATING CONDUCTIVITY VESSELS

κ_{18} for the purest water in a vacuum = $.04 \times 10^{-6}$ ohms⁻¹ cm.⁻¹ (Kohlrausch and Heydweiller); κ_{18} for conductivity water in air is about 10^{-6} ohms⁻¹ cm.⁻¹; KCl 1 n = normal KCl = 74.59 gm./litre at 18° C.; NaCl sat. = saturated NaCl at temp. t . of experiment. Unit—ohm⁻¹ cm.⁻¹. (See Kohlrausch, Holborn, and Dieselselhorst.)

Solution.	0° C.	8°	12°	16°	20°	24°
NaCl, sat..	'1345	'1688	'1872	'2063	'2260	'2462
KCl, 1 n .	'06541	'07954	'08689	'09441	'10207	'10984
KCl, 1/10 n .	'00715	'00888	'00979	'01072	'01167	'01264
KCl, 1/50 n .	'00152	'00190	'00209	'00229	'00250	'00271
KCl, 1/100 n .	'00078	'00097	'00107	'001173	'001278	'001386

87
CONDUCTIVITY OF SOLUTIONS

EQUIVALENT ELECTRIC CONDUCTIVITY Λ OF DILUTE AQUEOUS SOLUTIONS

Extrapolated numbers are indicated by (). Λ for infinite dilution is given under "O." Observers: inorganic solutes, Kohlrausch; organic, Bredig, *Zeit. Phys. Chem.*, 1894.

Solute at 18° C.	Gm. equiv. per litre = 1000 η .				Solute at 18° C.	Gm. equiv. per litre = 1000 η .			
	0	·0001	·01	·5		·0001	·0002	·01	·5
KCl . . .	130·1	129·1	122	102	$\frac{1}{2}$ CaCl ₂ . .	115·2	114·5	103	74·9
KBr . . .	132·3	131·1	124	105	$\frac{1}{2}$ SrNO ₃ . .	111·7	111·1	99	62·7
KI . . .	131·0	129·8	123	106	$\frac{1}{2}$ BaCl ₂ . .	[117/·0005]		107	77·3
KF . . .	111·3	110·5	104	83	$\frac{1}{2}$ MgCl ₂ . .	109·4	108·9	98·1	69·5
KSCN . .	121·3	120·2	114	95·7	$\frac{1}{2}$ ZnSO ₄ . .	109·5	107·5	72·8	—
KNO ₃ . .	126·5	125·5	118	89·2	$\frac{1}{2}$ CdNO ₃ . .	[100/·005]		96	63·9
NaCl . .	109·0	108·1	102	80·9	$\frac{1}{2}$ CuSO ₄ . .	109·9	107·9	71·7	—
NaF . .	90·15	89·3	83·5	60·0	$\frac{1}{2}$ PbN ₂ O ₆ . .	120·7	119·9	103	53·2
NaNO ₃ . .	105·3	104·5	98·2	74·0					
LiCl . .	98·9	98·1	99·2	70·7					
AgNO ₃ . .	115·8	115·0	108	77·5					
CsCl . .	133·6	132·3	125	—					
RbCl . .	—	132·3	125	—					
NH ₄ Cl . .	—	129·2	122	101					
TiCl . .	131·5	130·3	120	—					
					Acids.				
					HCl . . .	(377)	376	370	327
					HNO ₃ . .	(375)	374	368	324
					$\frac{1}{2}$ H ₂ SO ₄ . .	361	351	308	205
					$\frac{1}{3}$ H ₃ PO ₄ . .	(106)	102	85	—
					Alkalies.				
					KOH . . .	(234)	(233)	228	197
					NaOH . .	—	204·5	203·4	174
					NH ₃ . . .	53/·0002	38/·0005	9·6	1·35

Solute at 25° C.	Λ_{1024}	Λ_{∞}	Solute at 25° C.	Λ_{1024}	Λ_{∞}
Na formate . . .	98·1	100·4	Hydrochloride of—		
Na acetate . . .	85·7	87·5	-Propylamine . .	107·5	110·3
Na propionate . .	81·0	83·5	(CH ₃) ₄ PCl . . .	107·4	109·8
Na butyrate . . .	77·4	79·9	(C ₂ H ₅) ₄ PCl . . .	98·3	100·8
Na isobutyrate . .	77·7	80·1	(CH ₃) ₄ AsCl . . .	105·5	108·2
Hydrochlorides of—					
-Methylamine . .	125·1	127·8	Hydrochlorides of—	Λ_{256}	
-Ethylamine . . .	114·3	117·0	-Aniline	100·3	106·1
-Dimethylamine . .	117·5	120·3	-Methylamine . .	99·4	105·2
-Allylamine . . .	109·2	111·7	-o-Toluidine . .	97·4	103·7

EQUIVALENT ELECTRIC CONDUCTIVITY OF NON-AQUEOUS SOLUTIONS

$v = 1/m$ = volume in litres in which 1 gm. equivalent is dissolved. (See Tower, "Conductivity of Liquids," 1908.)

Solvent.	Solute.	t° C.	v	Λ	v	Λ	Solvent.	Solute.	t° C.	v	Λ	v	Λ
NH ₃	KBr	-38°	5740	317·6	12410	329·7	POCl ₃	N(C ₂ H ₅) ₄ I	25°	750	38·5	1500	44·3
"	AgNO ₃	-15	94	188	192	110	Formic acid	{ KCl	25	256	58	512	61
H ₂ CN	KI	0	392	298	1024	308	{ HCl	25	5·86	32·8	46·9	33·2	
"	S(CH ₃) ₃ I	0	512	327	1024	332	Acetone	KI	18	1157	155	2315	163
SO ₂	KI	0	1024	112·5	2048	134·5	"	LiCl	18	10	49·8	13·8	99·5
"	N(C ₂ H ₅) ₄ I	0	512	157·1	1024	167·7	"	AgNO ₃	18	288	15·7	576	17·6
AsCl ₃	N(C ₂ H ₅) ₄ I	25	150	63·2	750	59·7							

IONIC MOBILITIES

MOBILITIES OF IONS IN LIQUIDS

The mobility of the anion = $u_- = 1.037 \times 10^{-5} \Delta n$. (n = Hittorf's number.)
Example.—For KCl, $\Delta_\infty = 130.1$, $n = .505$, $\therefore u_- = 1.037 \times 10^{-5} \times .505 \times 130.1 = 6.8 \times 10^{-4}$ cm./sec. for Cl ions at 18° . Observers, Kohlrausch and Bredig; the latter's values have been multiplied by 1.1×10^{-5} to bring them to cm./sec.
Unit— 10^{-5} cm./sec. * $\frac{1}{2}$ Ca, etc.: the actual ionic velocity of the divalent ions is half the value stated here; these values, however, fit the equations given on p. 85.

Ion.	u 18°	Ion.	u 18°	Ion.	u 18°	Ion.	u 18°	Ion.	u 25°	Ion.	u 25°
H.	330	NH ₄	66.3	Zn*	48.4	F	48.3	HCO ₂	56.3	C ₂ H ₅ H ₃ N	51.5
Li	34.6	Tl	68.4	Cu*	49	Cl	67.8	CH ₃ CO ₂	42.1	(C ₂ H ₅) ₄ P.	33.7
Na	45.2	Ca*	53.7	Ag	56	Br	70	C ₂ H ₅ CO ₂	37.7	C ₆ H ₅ H ₃ N	39.5
K	67	Sr*	53.6	Cd*	49.2	I	68.8	n.C ₃ H ₇ CO ₂	33.8	aniline	39.5
Rb	70.5	Ba*	57.5	Pb*	63.5	NO ₃	64	Iso-	34.0	C ₆ H ₅ HN	48.5
Cs	70.5	Mg*	47.7	OH	180	SO ₄ *	71	CH ₃ H ₃ N	53.4	(CH ₃) ₄ As.	41.8

DIRECTLY OBSERVED MOBILITIES

Deduced from the observed movement of an ionic boundary. m = equivalent concentration. **Unit**— 10^{-5} cm./sec. at 18° C. (See Denison and Steel, *Phil. Trans.*, 1906.)

Ion.	m	u	Ion.	m	u	Ion.	m	u	Ion.	m	u	Ion.	m	u
K	.5	55.3	Na	.1	31.8	Ba	.5	33	Mg	.2	16.7	Cl	.5	52.9
												SO ₄	.2	30.4

ELECTROMOTIVE FORCES AND RESISTANCES OF CELLS

The E.M.F.'s given are for cells on open circuit, and are only approximate; in the case of primary batteries they refer to freshly made up cells. The internal resistances quoted are only typical; they vary very widely in practice. With many primary cells the E.M.F. drops and the internal resistance increases as the cell ages. Nearly all modern dry cells are modified Leclanché batteries.

(See Slingo and Brooker's "Electrical Engineering.")

Cell.	Description.	E.M.F.	Resistance.
Bichromate . . .	Zn and C in 1 vol. strong H ₂ SO ₄ and 20 vols. sat. K ₂ Cr ₂ O ₇ sol.	Volts. c. 2.0	Ohms. very low
Bunsen . . .	Zn in 1 vol. H ₂ SO ₄ and 12 vols. H ₂ O; C in strong HNO ₃	1.8-1.9	—
Clark (see p. 8)	Zn amalgam and Hg in sat. ZnSO ₄ sol.	1.433	c. 500
Daniell . . .	Zn in ZnSO ₄ sol. or H ₂ SO ₄ (1 to 12); Cu in sat. CuSO ₄ sol.	1.07-1.08	c. 4
Grove . . .	Like Bunsen with Pt instead of C	1.8-1.9	—
Leclanché . . .	Zn and C in NH ₄ Cl, C, and MnO ₂	c. 1.5	0.25-4
Secondary . . .	Pb and PbO ₂ (etc.) in H ₂ SO ₄ of density 1.2	2.2-1.9	negligible
Tucker . . .	"Hygroscopic cell." Zn and C with sat. CaCl ₂ sol.	1.4	—
Weston (see p. 8).	Cd amalgam and Hg in sat. CdSO ₄ sol.	1.018	c. 500

MAGNETIC INDUCTION

 \mathcal{H} = magnetic force \mathcal{I} = intensity of magnetization= magnetic moment per cm.³= pole strength per cm.² \mathcal{B} = magnetic induction, or flux density= $\mathcal{H} + 4\pi\mathcal{I}$ μ = permeability = \mathcal{B}/\mathcal{H} . See p. 6. H = susceptibility = $\mathcal{I}/\mathcal{H} = (\mu - 1)/(4\pi)$. See p. 6.**Coercivity**, $\mathcal{H}_B = 0$, is the demagnetizing force required to make $\mathcal{B} = 0$ after saturation.**Coercive force** is the demagnetizing force required to make $\mathcal{B} = 0$ after some particular field strength.**Remanence**, $\mathcal{B}_R = 0$, is the induction remaining when the magnetic force is removed after some particular field strength.

The work done, *i.e.* **hysteresis loss**, Q_h , in taking a cm.³ of magnetic material through a magnetic cycle between limits $\pm H_m = \int \mathcal{H} d\mathcal{B} = \frac{1}{4\pi} \int \mathcal{H} d\mathcal{B}$. Steinmetz's empirical formula for the hysteresis loss is $\eta \mathcal{B}_{\max}^n$, where η is a constant, and generally $n = 1.6$. The magnetic properties of a material depend not only on its chemical composition, but on its previous mechanical and heat treatment; thus only general characteristics are indicated below.

Heusler alloys (discovered by Heusler in 1903) are composed of Cu, Mn, and Al. They do not show the Kerr effect.Good permanent **magnet steel** contains about 5% W and 6% C, is free from Mn, Cu, Ni, and Ti, and is hardened at 850° C. (Hannack, 1909). Cast iron, chilled from 1000° C., may also be used (Peirce and Campbell).

References.—Pure iron, Peirce, *Amer. Jour. Sci.*, 27 and 28, 1909; Terry, *Phy. Rev.*, 1909; iron and manganese, Burgess and Aston, *Phil. Mag.*, 1909; Heusler alloys, Stephenson, *Phy. Rev.*, 1910. (Ewing, "Magnetic Induction in Iron," and Kohlrausch, "Prakt. Phys.")

Material.	Permeability μ .						Coer- civity.	Rema- nence.	\mathcal{H}_c .	Hyst. loss, Q_h .
	$\mathcal{H} = 5$	$\mathcal{H} = 1$	$\mathcal{H} = 5$	$\mathcal{H} = 20$	$\mathcal{H} = 60$	$\mathcal{H} = 150$				
Swedish wrought iron	2500	3710	2060	736	274	120	0.8	4,000	200	ergs/cm. ³ 6,700
Annealed cast steel	1450	3500	2100	747	280	123	0.97	7,100	151	11,700
Unannealed cast steel	490	970	1700	680	270	122	2.08	9,000	156	20,400
Cast iron	—	—	81	182	117	65	11.9	4,230	155	34,300
Magnet (Hardened steel)	—	—	68	15	78	193	52.6	11,700	234	211,000
Tungsten	—	—	80	10	119	204	105	9,880	505	116,000

Material.	\mathcal{H}_{\max} .	Induction, \mathcal{B} , for		μ_{\max} .	For \mathcal{H}_{\max} .		
		\mathcal{H}_{\max} .	$\mathcal{H} = 100$.		Coer.	Reman.	Hyst. loss.
Mild steel**	129	18,190	17,700	8350	0.6	10,300	ergs/cm. ³ 4,900
Steel, 2.8% Cr, .8% C.	—	—	—	—	56	6,400	—
" 5.5% W, .6% C.	Hardened at 770°	—	—	—	72	7,000	280,000
" 7.7% W, 1.9% C.	" 800°	—	—	—	85	4,700	—
" 4% Mo, 1.2% C.	" 800°	—	—	—	85	6,700	—
Iron†	50	17,100	—	1750	2.2*	c. 53% \mathcal{B}_{\max}	—
Silicon iron, .6% Si†	50	16,000	—	1900	1.6*	c. 43% "	—
" .45% Si†	50	15,100	—	2500	1.2*	c. 39% "	—
Electrolytic iron (very pure)§	210	21,250	—	—	18	10,000	—
	Heated to 1200° C.	16,000	—	—	2.5	12,500	—
Hadfield's manganese steel	—	—	—	13-15	—	v. small	—
Nickel, annealed	100	5,137	—	296	8	3,570	—
Cobalt	140	10,000	9,500	174	12	3,400	—
" .96%	114	8,237	7,800	177	—	—	19,000
Heusler alloy¶	92	2,735	—	115	—	—	—

* $H = 10$.† Otto, *Deut. Phys. Ges. Berlin*, 1910.

‡ Bar magnet.

§ Burgess and Taylor, 1906.

|| .12% Mn, 1% C.

¶ 24 Mn, 16 Al, 60 Cu. McClellan, 1907.

** Gumlich and Schmidt (Reichsanstalt), 1901.

MAGNETISM

MAGNETIC SUSCEPTIBILITIES OF THE ELEMENTS, ETC.

The susceptibility $H = \frac{I}{H} = (\mu - 1)/(4\pi)$. $H = 0$ for a vacuum. The susceptibility depends very much on the purity of the material, especially upon the absence of iron. For pure elements H appears to be independent of μ , except possibly in the case of Mg, Sb, and Ru. H is a periodic property of the atomic weight; for example, P, As, Sb, and Bi are comparatively strongly diamagnetic.

The values below are per grm. at 18°C ., except where some temperature is specified. The gases are per cm^3 at 1 atmos. [Honda (*Ann. d. Phys.*, 1910) used purest available materials and corrected H for any traces of iron; see also P. Curie, *Œuvres*, Paris, 1908.] + means paramagnetic; -, diamagnetic.

Elem.	H	Obs.	Elem.	H	Obs.	Elem.	H	Obs.
Solids			Solids			Solids		
Al	$\times 10^{-6}$ + '65	L., W., H.	(contd.) $\times 10^{-6}$			(contd.) $\times 10^{-6}$		
Sb	- '95	H.	P.	- '9	H., B., C., Q.	V	+ '15	H.
As	- '31	H.	Pt.	+ '132	—	Zn	- '15	K., L., H.
Bi	- '14	B. C. D. E. W.	K.	+ '4	H.	Zr	- '45	H.
B.	- '71	H.	Rh.	+ '11	H., F.	Liquids		
Cd	- '17	H.	Ru.	+ '56	H.	Br	- '41	C., Q.
Cr	+ '37	H.	Se.	- '32	H., C.	Hg	- '19	Q., M., H.
Cu	- '087	H.	Si.	- '12	H.	N liq.	+ '28	F., D.
Au	- '15	K., H.	Ag.	- '2	H.	O liq.	+ 324	F., D.
I.	- '36	B., C., H.	Na.	+ '51	H.	$\text{H}_2\text{O}, 15^\circ$	- '837	Du B.
Ir.	+ '15	H.	S.	- '5	B., C., L., K., H.	$\text{H}_2\text{O}, 15^\circ$	- '77	S.
Fe	See p. 89.		Ta.	+ '93	H.	Gases		
Pb	- '12	H., K., L.	Te.	- '32	E., C., H.	Air, 16°	+ '032	Du B.
Mg	+ '55	H.	Tl.	- '3	H.	A.	- '010	T.
Mn	+ '106?	H.	Th.	+ '18	H.	He	- '002	T.
Mo	+ '04	H.	Sn.	+ '025	K., H.	H	- '008	Q.
Nb	+ '13?	H.	Ti.	- '2	H.	N	+ '024	Du B.
Os	+ '04	H.	W.	+ '33	H.	O	+ '123	Du B., Q.
Pd	+ '58	H., K., C., F.	U.	- '9	M., H.			

B., E. Becquerel, 1855; C., Curie, 1895; D., Dewar, 1892; Du B., Du Bois; E., Ettingshausen; F., Finke; F. D., Fleming and Dewar; H., Honda; K., Königsberger, 1901; L., Lombardi, 1897; M., St. Meyer; Q., Quincke; S., Scarpa, 1905; T., Tänzler, 1907; W., Wills, 1898.

TEMPERATURE AND MAGNETIZATION

The magnetic moment (M) of a magnet diminishes as the temperature (t) rises. In $M_t = M_0(1 - \alpha t)$, α varies widely, but is of the order '0003 to '001. The permeability μ also depends on the temperature. There is a **critical temperature** above which μ is very small; in the case of iron it is one of the recalcrescence temperatures, and is the same as for carbon steels containing up to 45 % of C.

The critical temperature of a metal is not perfectly definite, but depends to some extent on whether the metal is being heated or cooled.

Substance.	Crit. Temp.	Observer.	Substance.	Crit. Temp.	Observer.
Iron	690°-870° C.	Hopkinson	Nickel, 95% .	310°	Hopkinson
"	c. 895	Roberts-Austen	"	377	Weiss, 1907
"	855-867	Osmond	Magnetite . .	582	"
"	757	Weiss, 1907	Heusler alloys	c. 300	Gray, 1908
Cobalt . . .	1075	Stifler, 1911	Stalloy . . .	760	Hadfield

Nickel steel (25 % Ni); 0° to $50^\circ \mu = 1.4$ to 60; 50° to $580^\circ \mu = 60$ to 0.4.

STEINMETZ'S COEFFICIENT

Values of η in Steinmetz's formula $\eta B_{\max}^{1.6}$ for the hysteresis loss in ergs per c.c. per cycle. B_{\max} is the maximum value of the induction.

Substance.	η	Substance.	η
3½% Silicon iron (Stalloy) .	·0007	Grey cast iron	·013
Good transformer iron . . .	·0011	Nickel	·012 to ·038
Dynamo cast steel	·0026	Cobalt	·012
High carbon steel, hardened	·025		

TERRESTRIAL MAGNETIC CONSTANTS

Magnetic observatories no longer remain in large cities owing to electric tram disturbances, and thus many of the places for which reliable data exist are not generally known. The general locality of the station is indicated in many cases below.

Magnetic constants obtained in most physical laboratories are usually abnormal owing to the proximity of iron in some form.

Much of the data below is derived from the Reports of Kew Observatory, and the publications of the United States Coast and Geodetic Survey.

A W declination means that the N-seeking end of the magnetic needle points west of true north; a N inclination means that the same end of the needle points downwards.

H and V are the horizontal and vertical components of the earth's magnetic field.

(See Chree, "Terrestrial Magnetism," Encyc. Brit., 11th edit., 1911.)

Place.	Latitude.	Longi- tude.	Year.	Declina- tion.	Inclina- tion.	H.	V.
	° ' "	° ' "		° ' "	° ' "	c.g.s.	c.g.s.
North magnetic pole . . .	70 5 N	96 45 W	—	—	90 0 N	0	—
South magnetic pole* . .	72 25 S	154 E	1908	—	90 0 S	0	—
British Isles—							
Aberdeen (University) .	57 9 N	2 7 W	1909	16 34 W	70 39 N	·163	·464
Eskdalemuir (Dumfries) .	55 19 N	3 12 W	1909	18 30 W	69 39 N	·1684	·4519
Falmouth (Cornwall) . .	50 9 N	5 5 W	1909	17 48 W	66 31 N	·1880	·4327
Greenwich	51 28 N	0 0	1916	14 47 W	66 53 N	·1849	·4333
Kew	51 28 N	0 19 W	1909	16 11 W	67 0 N	·1851	·4359
Leeds (University) . . .	53 48 N	1 33 W	1909	18 2 W†	68 35 N	·176	·449
St. Helier (Jersey) . . .	49 12 N	2 5 W	1907	16 27 W	65 35 N	—	—
Stonyhurst (Lancs.) . .	53 51 N	2 28 W	1909	17 29 W	68 43 N	·1742	·4472
Valencia (S. W. Ireland)	51 56 N	10 15 W	1909	20 50 W	68 15 N	·1788	·4481
Africa—							
Cape Town	33 56 S	18 29 E	1885	30 15 W	56 0 S	·199	·295
Helwan (Cairo)	29 52 N	31 21 E	1908	2 56 W	40 39 N	·3003	·2579
Mauritius	20 6 S	57 33 E	1908	9 14 W	53 45 S	·2342	·3193
America—							
Agincourt (Toronto) . .	43 47 N	79 16 W	1906	5 45 W	74 36 N	·1640	·5950
Cheltenham (Washing- ton)	38 44 N	76 50 W	1909	5 34 W	70 31 N	·1988	·5620
Fairhaven (Mass.) . . .	41 37 N	70 54 W	1908	12 27 W	73 8 N	·1736	·5724
Goat Island (California)	37 49 N	122 22 W	1909	17 53 E	62 11 N	·2525	·4786
Greenwich (New York) .	41 0 N	73 37 W	1908	10 14 W	72 13 N	·1822	·5680
Rio de Janeiro	22 55 S	43 11 W	1906	8 55 W	13 57 S	·2477	·0616
Santiago (Chili)	33 27 S	70 42 W	1906	14 19 E	30 12 S	—	—
Sitka (Alaska)	57 3 N	135 20 W	1909	30 12 E	74 37 N	·1557	·5659
Waukegan (Chicago) . .	42 21 N	87 51 W	1908	2 39 W	72 46 N	·1830	·5898

* Mawson and David (with Shackleton), 1908.

† 1907.

TERRESTRIAL MAGNETISM

TERRESTRIAL MAGNETIC CONSTANTS (contd.)

Place.	Latitude.	Longitude.	Year.	Declination.	Inclination.	H.	V.
							c.g.s.
Asia—							
Alibag (Bombay)	18 39 N	72 52 E	1908	1 2 E.	23 22 N	3686	1592
Barrackpore (Calcutta) .	22 46 N	88 22 E	1907	1 10 E	30 30 N	3729	2197
Hong Kong	22 18 N	114 10 E	1909	0 2 E	31 1 N	3709	2239
Australasia—							
Christchurch (N.Z.) . . .	43 32 S	172 37 E	1903	16 18 E	67 42 S	2266	5526
Honolulu (Hawaii) . . .	21 49 N	158 4 W	1909	9 26 E	40 54 N	2917	2527
Melbourne	37 59 S	144 58 E	1901	8 27 E	67 25 S	2331	5602
Sydney	33 52 S	151 12 E	1885	9 30 E	62 30 S	268	515
Europe—							
Arctic (Norway)	69 56 N	22 58 E	1903	0 43 W	76 21 N	1258	5178
Regions (Spitzbergen). .	77 41 N	14 50 E	1903	10 55 W	80 8 N	0942	5417
Odessa	46 24 N	30 48 E	1901	4 27 W	62 18 N	2188	4168
Pawlowsk (Petrograd) . .	59 41 N	30 29 E	1906	1 4 E	70 37 N	1653	4696
Potsdam	52 23 N	13 4 E	1909	9 11 W	66 20 N	1883	4297
Rude Skov (Copenhagen)	55 51 N	12 27 E	1908	9 43 W	68 45 N	1741	4476
Uccle (Brussels)	50 48 N	4 21 E	1908	13 37 W	66 2 N	1906	4287
Val Joyeux (Paris) . . .	48 49 N	2 1 E	1909	14 33 W	64 44 N	1973	4179

SECULAR MAGNETIC CHANGES

At the present period we are going through a remarkable secular alteration. For generations H had been steadily rising in Western Europe, but during the last few years a wave of depression has travelled across from the east. H has steadily fallen at Petrograd since about 1900, at Potsdam since about 1905, at Greenwich and Kew since 1907, while in 1909 H was still rising at Falmouth and Valencia. The easterly motion of the declination needle has also increased notably since 1900. Thus secular change data based on, say, the last five years will not serve to prospect the future.

Mean change per annum at	1908-1909.		1904-1909.			
	Decln.	H.	Decln.	Incln.	H.	V.
Greenwich	-5.9	-5 × 10 ⁻⁵	-5.5	-0.7	+1 × 10 ⁻⁵	-20 × 10 ⁻⁵
Kew	-6.1	-9 "	-5.4	-1.1	+2 "	-35 "
Stonyhurst	-7.0	-10 "	-5.9	-1.1	+6 "	-25 "
Falmouth	-6.3	+4 "	-4.7	-1.4	+9 "	-30 "
Valencia	-5.4	+7 "	-5.0	-1.2	+7 "	-25 "

SECULAR CHANGES AT LONDON (GREENWICH)

Year.	Decln.	Incln.	Year.	Decln.	Incln.	H.
1580	11 17 E	72 0 N	1851	22 25 W	68 47 N	1729
1660	0 0	73 15 N	1875	19 21 W	67 42 N	1795
1720	13 0 W	74 40 N*	1907	16 0 W	66 56 N	1853*
1815	24 27 W*	70 30 N	1916	14 47 W	66 53 N	1849

* Maximum.

SPARKING POTENTIALS

The sparking voltages given below are those which will break down non-ionized air at atmospheric pressure and room temperature. The electrodes are equal smooth polished metal balls of various diameters. Russell (*Phil. Mag.*, 1906) gives the dielectric strength of air at atmospheric pressures as between 38,000 and 39,000 volts per cm. for either direct or alternating potentials.

(See Kaye's "X Rays" (Longmans, 1916) for further values.)

Spark gap.	Diameter of balls in cms.				Spark gap.	Diameter of balls in cms.			
	0.5	1.0	2.0	5.0		0.5	1.0	2.0	5.0
cm.	volts. $\times 10^3$	volts. $\times 10^3$	volts. $\times 10^3$	volts. $\times 10^3$	cm.	volts. $\times 10^3$	volts. $\times 10^3$	volts. $\times 10^3$	volts. $\times 10^3$
0.1	4.8	4.8	4.7	—	0.9	19.6	25.6	28.6	30.1
0.2	8.4	8.4	8.1	—	1.0	20.2	26.7	30.8	32.7
0.3	11.3	11.4	11.4	—	1.5	22	31.6	39	46
0.4	13.8	14.4	14.5	—	2.0	23	36	47	58
0.5	15.7	17.3	17.5	18.4	3.0	24	42	57	77
0.6	17.2	19.9	20.4	21.6	4.0	25	45	64	92
0.7	18.3	22.0	23.2	24.6	5.0	26	47	69	105
0.8	19.0	24.1	26.0	27.4					

HOMOGENEOUS X-RAYS

Mass absorption coefficients, λ/ρ , measured in Al foil. λ is the absorption coefficient (see p. 107) of the homogeneous characteristic (K) X radiation from a metal; ρ is the density of aluminium foil. For a complete set of values, see Kaye's "X Rays" (Longmans, 1916).

Radiator.	Al	Cr	Fe	Ni	Co	Cu	Zn	As	Se	Ag
λ/ρ	3400	136	88.5	59.1	71.6	47.7	39.4	22.5	18.5	2.5

CATHODE DARK SPACE

The thickness (d) of the Crookes dark space is given by $d = (A/p) + B/\sqrt{i}$, where p is the pressure, i the current density, and A and B are constants for each gas. This equation is satisfied very exactly by the ordinary elementary gases, and a little less so by the gases of the helium group. Unfortunately for the use of the dark space as a pressure indicator, the current density term in the formula is almost as large as the pressure term for pressures about 1/10 mm.

The values of A and B below are for large plane aluminium electrodes. d is measured in cms., p in mms. of mercury. The unit of i is 1/10 milliampere per sq. cm. of cathode, which is about the sort of current density that obtains with an average coil discharge and a moderate-sized cathode.

(See Aston and Watson, *Proc. Roy. Soc.*, 1911.)

Gas.	Hydrogen	Nitrogen	Air	Oxygen
A	.26	.068	.065	.057
B	.43	.40	.42	.50

RECOMBINATION AND DIFFUSION

COEFFICIENTS OF RECOMBINATION α

α is given below in terms of $1000e$, where e is the numerical value of the ionic charge: 4.7×10^{-10} in electrostatic units. For air, $\alpha = 3320e = 1.56 \times 10^{-10} \text{ cm.}^3 \text{ sec.}^{-1}$. Room temp. and pressure.

Gas.	Air.	O ₂	CO ₂	H ₂
α	3.42, T.; 3.38, Mc.; 3.2, L.; 3.3, H.; 3.32*, E.	3.38, T.	3.5, T.	3.02, T.; 2.94, Mc.

E., Erikson, *P.M.*, 1909; H., Hendren, *P.R.*, 1905; L., Langevin, *A.C.P.*, 1902; Mc., McClung, *P.M.*, 1902; T., Townsend, *P.T.*, 1899. * 17° C., 760 mm. Hg.

 α IN AIR AND PRESSURE

Press. in atmos.2	.5	1	2	3	5	L., Langevin. H., Hendren.		
α (relative values), L.	5	12	27	30	26	18			
Press. in cms.	76	45	25	15	10	5	3.5	2	1
α (absolute values), H.	3.3	2.65	2.07	1.75	1.55	1.31	1.25	1.15	1.00

 α IN AIR AND TEMPERATURE

Air at constant density. (E., Erikson; P., Phillips, *Electrician*, 1909.)

Temp. °C.	-179	-68	12	64	100	155	Temp. °C.	15	100	155	176
α (in terms $1000e$), E.	7.5	5.6	3.47	2.31	1.73	1.38	α (relative values), P.	1	.50	.40	.36

IONIC COEFFICIENTS OF DIFFUSION D

Rate of interdiffusion (in $\text{cm.}^3 \text{ sec.}^{-1}$) of gaseous ions in dry air: D_+ for positive, D_- for negative ions. (Townsend, *Phil. Trans.*, 1899, 1900.)

Ionisation	Röntgen Rays.	β and γ Rays.	Ultra-violet light.	Point discharge.
D_+ at 76 cm.028	.032	—	.0247, .0216
D_- at 76 cm.043	.043	.043	.037, .032

GASES IONIZED BY RÖNTGEN RAYS

Air, CO₂, and hydrogen at 15° C. and 760 mm.

Dry Gas.	D ₊	D ₋	Dry Gas.	D ₊	D ₋	Moist Gas.	D ₊	D ₋	Moist Gas.	D ₊	D ₋
Air {dried by CaCl ₂ }	.028	.043	CO ₂ {dried by CaCl ₂ }	.023	.026	Air {sat. with H ₂ O}	.032	.035	CO ₂ {sat. with H ₂ O}	.024	.025
O ₂ {dried by CaCl ₂ }	.025	.04	H ₂ {dried by CaCl ₂ }	.123	.19	O ₂ {sat. with H ₂ O}	.029	.036	H ₂ {sat. with H ₂ O}	.128	.142

AIR IONIZED BY β AND γ RAYS

Press. p. in cms.	77.2	55	40	30	20	Press. p. in cms.	77.2	55	40	30	20
D_+ at 15° C.	.0317	.042	.0578	.078	.118	D_- at 15° C.	.0429	.0542	.078	.103	.155
pD_+ „	2.45	2.31	2.31	2.34	2.36	pD_- „	3.3	2.98	3.12	3.09	3.1

A.C.P., *Ann. de Chim. et de Phys.*; *P.M.*, *Phil. Mag.*; *P.R.*, *Physical Review*; *P.T.*, *Phil. Trans.*

MOBILITIES OF IONS IN GASES

Velocities of ions are in cm. per sec. for unit field, or in $\text{cm.}^2 \text{ sec.}^{-1} \text{ volt}^{-1}$ at temp. and press. of room. K_+ = mobility of positive ion, K_- of negative.

For **moist air** (i.e. saturated with H_2O), $K_+ = 1.37$, $K_- = 1.51$.

For **dry air** (dried by CaCl_2), $K_+ = 1.36$, $K_- = 1.87$. (Zeleny (air blast method), *Phil. Trans.*, 1900.) * Mean = $(K_+ + K_-)/2$.

For mobilities of natural ions in air, see p. 105.

Dry Gas.	K_+	K_-	Ionization and Observer.	Dry Gas.	K_+	K_-	Ionization and Observer.
	76 cm. Hg				76 cm. Hg		
Air	1.32	1.80	Point disch., Chattock, <i>P.M.</i> , 1899, 1901.	CO ₂	0.76	0.81	X-rays, Zeleny, 1900.
"	1.54	1.78	X-rays, Wellisch, <i>Phil</i> <i>Trans.</i> , 1909.	"	0.86	0.90	" Langevin, '03.
"	1.40	1.70	" Langevin, <i>A.C.P.</i> , 1903.	HCl	0.81	0.85	" Wellisch, '09.
"	1.39	1.78	" Phillips, <i>P.R.S.</i> , 1906.	SO ₂	1.27*	0.41	" Rutherford.
"	1.36	1.87	" Zeleny, <i>Phil.</i> <i>Trans.</i> , 1900.	Cl ₂	0.44	0.41	" Wellisch, '09.
"	1.40	1.78	Mean value.	N ₂ O	1.0*	0.90	" Rutherford.
H ₂	5.4	7.43	Point disch., Chattock.	NH ₃	0.82	0.90	" Wellisch, '09.
"	6.7	7.9	X-rays, Zeleny, 1900.	Me. acetate	0.74	0.80	" "
He	5.09	6.31	" Franck and Pohl, <i>V.D.P.G.</i> , '07.	Me. bromide	0.33	0.36	" "
N ₂	1.6*	—	X-rays, Rutherford, <i>P.M.</i> , 1897.	Me. iodide	0.29	0.28	" "
O ₂	1.36	1.80	" Zeleny, 1900.	Et. alcohol	0.21	0.22	" "
"	1.3	1.85	Point disch., Chattock.	Et. acetate	0.34	0.27	" "
CO	1.1	1.14	X-rays, Wellisch, '09.	Et. aldehyde	0.31	0.28	" "
CO ₂	0.83	0.92	Point disch., Chattock.	Et. ether	0.31	0.30	" "
				Et. chloride	0.33	0.31	" "
				Et. ether	0.29	0.31	" "
				Et. formate	0.30	0.31	" "
				Et. iodide	0.17	0.16	" "
				C.Cl ₄	0.30	0.31	" "
				Pentane	0.36	0.35	" "
				Acetone	0.31	0.29	" "

IONIC MOBILITY AND PRESSURE

Air ionized by Röntgen rays. (Langevin, *A.C.P.*, 1903.)

Press. cm.	7.5	20	41.5	76	143.5	Press. cm.	7.5	20	41.5	76	142
K_+	14.8	5.45	2.61	1.40	0.75	K_-	21.9	7.35	3.31	1.7	0.9

IONIC MOBILITY AND TEMPERATURE

Air at 76 cm. press. ionized by Röntgen rays. (Phillips, *P.R.S.*, 1906.)

Temp. ° C.	138°	126°	110°	100°	75°	60°	12°	-64°	-179°
K_+	2.00	1.95	1.85	1.81	1.67	1.60	1.39	0.945	0.235
K_-	2.49	2.40	2.30	2.21	2.12	2.00	1.785	1.23	0.235

IONIC MOBILITIES IN LIQUIDS AND SOLIDS

Ionized by radium rays. (Bohm-Wendt and v. Schweidler, *Phys. Zeit.*, 1909; Bialobjeski, *Compt. Rend.*, 1909.)

Substance.	$(K_+ + K_-)$	Substance.	$(K_+ + K_-)$
Petroleum ether	3.8×10^{-4}	Ozokerite at 100°	5.1×10^{-4}
Vaseline	5.3×10^{-6}	" " 80°	35.0×10^{-4}

A.C.P., *Ann. de Chim. et de Phys.*, *P.M.*, *Phil. Mag.*; *P.R.S.*, *Proc. Roy. Soc.*; *V.D.P.G.*, *Verh. Deutsch. Phys. Gesell.*

K in cm. sec.⁻¹ per volt cm.⁻¹ for coal-gas flames in most instances. The ionic mobility is independent of the acid of the salt. Gold's and Wilson's values for K agree the best with existing theory, which makes $K = X e \lambda / m u = 17,000$ at 1800° C. (Gold). X is the electric field per cm., λ is the mean free path, and u the velocity of the corpuscle.

Salt.	Temp.	K ₊	K ₋	Observer.
Cs, Rb, K, Na, Li . .	Flame, <i>c.</i> 2000° C.	62	<i>c.</i> 1000	H. A. Wilson, <i>P.T.</i> , 1899
1/20 normal KCl . . .	Flame	260	1400	Marx. <i>Ann. der Phys.</i> , 1900
NaCl	"	340	1800	
1/256 normal K salt . .	Flame, <i>c.</i> 2000°	—	1320	Moreau, <i>Journ. de Phys.</i> , 1903
1/16 normal Na salt . .	" "	—	1280	
Concentrated sols. of alkalies	" "	80	—	
Cs, Rb, K, Na, Li . .	Air at 1000°	7·2	26	H. A. Wilson, <i>P.T.</i> , 1899 and <i>P.M.</i> , 1906
Ba, Sr, Ca	" "	3·8	—	
K, Na	Flame, <i>c.</i> 1800°	—	8000	Gold, <i>P.R.S.</i> , 1907, ratio of potential grad. to current
K	Flame, <i>c.</i> 1800°	—	13,000	Poten. grad., and gas velocity
K ₂ CO ₃	Bunsen burner	—	9600	H. A. Wilson, <i>P.R.S.</i> , 1909
Na	Flame, <i>c.</i> 2000°	—	1170	Moreau, <i>C.R.</i> , 1909

Expansion = v_2/v_1 , where v_1 is the volume of the gas before, and v_2 the volume after expansion. **Supersaturation** of the vapour (at end of cooling by expansion) necessary for condensation = $S = (\text{density of vapour when drops are formed})/(\text{density of saturated vapour at the same temp.})$. (See J. J. Thomson, "Conduction of Electricity through Gases.")

Dust-free gas saturated with water-vapour. (C. T. R. Wilson, *P.T.*, '97, '99, '00.)

Gas.	Rain-like Condensation.		Cloud-like Condensation.		Gas.	Rain-like Condensation.		Cloud-like Condensation.	
	v_2/v_1	S.	v_2/v_1	S.		v_2/v_1	S.	v_2/v_1	S.
Air . .	1·252	4·2	1·38	7·9	CO ₂ . .	1·365	4·2	1·535	7·3
O ₂ . .	1·257	4·3	1·38	7·9	Cl ₂ . .	1·3	3·4	1·45	5·9
N ₂ . .	1·262	4·4	1·38	7·9	H ₂ . .	—	—	1·38	7·9

(L., Laby, *Phil. Trans.*, 1908; P., Przibram, *Wien Ber.*, 1906.) •

Vapour and Observer.	Ion.	v_2/v_1	S.	Vapour and Observer.	Ion.	v_2/v_1	S.
Water (C. T. R. Wilson)	-	1·25	4·15	n-Butyric acid, L. . . . ?	?	1·38	150
Water (C. T. R. Wilson)	+	1·31	5·8	iso-Butyric acid, L. . . . ?	?	1·36	130
Et. acetate, L. . . . +	+	1·48	8·9	iso-Valeric acid, L. . . . ?	?	1·22	6'3
Me. butyrate, L. . . . +	+	1·33	5·3	Methyl alcohol, P. . . . +	+	1·25	3'1
Me. iso-butyrate, L. . . ?	?	1·35	5·2	Ethyl alcohol, P. . . . +	+	1·17	2'3
Propyl-acetate, L. . . +	+	1·31	5·0	Propyl alcohol, P. . . . ?	?	1·18	30
Et. propionate, L. . . ?	?	1·41	7·8	iso-Butyl alcohol, P. . . ?	?	1·2	3'6
Formic acid, L. . . . ?	?	1·78	25·1	iso-Amyl alcohol, P. . . +	+	1·22	5'5
Acetic acid, L. . . . +	+	1·44	9·3	" " L. . . . +	+	1·18	4'1
Propionic acid, L. . . ?	?	1·34	9·4	Chloroform, P. . . . +	+	1·54	3'0

A.C.P., Ann. de Chim. et de Phys.; C.R., Compt. Rend.; P.M., Phil. Mag.; P.R.S., Proc. Roy. Soc.; P.T., Phil. Trans.

NE FOR ELECTROLYTIC IONS

NE is given both in electrostatic units (E.S.U.) and electromagnetic units (E.M.U.).

N is the number of molecules in a c.c. of gas at 76 cm. Hg ($g = 980.6$) and $t^\circ \text{C.}$, and E is the charge on the monovalent ion in electrolysis.

Antecedent data.—1 coulomb deposits 1.11827 mgm. Ag. At. wt. of Ag, 107.88; of H, 1.008. Density of $\text{H}_2 = 8.987 \times 10^{-5}$ gm. per c.c. at 0°C.

Gas.	E.S.U.	E.M.U.	Gas.	E.S.U.	E.M.U.	Gas.	E.S.U.	E.M.U.
	$\times 10^{10}$			$\times 10^{10}$			$\times 10^{10}$	
H_2 at 0°C.	1.29015	0.4300	O_2 at 0°	1.2924	0.4308	Ideal gas at 0°	1.2913	0.43044
H_2 at 15°C.	1.2230	0.4077	O_2 at 15°	1.2248	0.4083	gas at 15°	1.2241	0.40803

Ne FOR GASEOUS IONS

N is the number of molecules per c.c. of **air** at room temp. and 76 cm. Hg; e is the ionic charge in E.S.U., e_- for negative and e_+ for positive ions.

Ionization.	Ne ₋	Ne ₊	Observer.
X rays	1.23×10^{10}	2.41×10^{10}	Townsend, <i>P.R.S.</i> , 1908, 1909.
Ra rays	1.24×10^{10}	1.26 to 1.37×10^{10}	Haselfoot, <i>P.R.S.</i> , 1909.

Ne CALCULATED

In E.S.U., $Ne = 3.04 \times 10^8 \times K/D = 3.04 \times 10^8 \times 1.40/0.028 = 1.52 \times 10^{10}$ for positive air ions at 76 cm. and room temp. For D and K, see pp. 94, 95.

Gas.	Ne ₊	Ne ₋	Gas.	Ne ₊	Ne ₋		Ne ₊	Ne ₋
Air .	$1.52 \cdot 10^{10}$	$1.26 \cdot 10^{10}$	H_2 .	$1.50 \cdot 10^{10}$	$1.23 \cdot 10^{10}$	Mean{	$1.42 \cdot 10^{10}$	$1.22 \cdot 10^{10}$
O_2 .	$1.62 \cdot 10^{10}$	$1.38 \cdot 10^{10}$	CO_2 .	$1.07 \cdot 10^{10}$	$1.02 \cdot 10^{10}$		$1.32 \cdot 10^{10}$	

THE IONIC CHARGE e

$e = 4.77 \times 10^{-10}$ E.S.U. = 1.59×10^{-20} E.M.U., as a mean of the latest determinations. See Millikan, *P.M.*, July, 1917.

Ionization.	Method.	e in E.S.U.	Observer.
Röntgen rays; negative ions.	By measuring total charge on a cloud and obtaining number of ions from size of drops by Stokes' law.	$6.5 \cdot 10^{-10}$	J. J. Thomson, <i>P.M.</i> , 1898.
Ultra-violet light on metal; negative ions.		6.8 "	J. J. Thomson, <i>P.M.</i> , 1899.
Röntgen rays; negative ions.	Force (by Stokes' law) exerted by an electric field on a singly charged drop.	3.1 "	H. A. Wilson, <i>P.M.</i> , 1903.
Radium rays; negative ions.	The observer's original method.	3.4 "	J. J. Thomson, <i>Proc. Camb. Phil. Soc.</i> , 1903.
Charged spray of electrolytic O_2 .	Total charge on a cloud. No. of ions from weight of cloud and size of drops, using Stokes' law.	3.0 "	Townsend, <i>Proc. Camb. Phil. Soc.</i> , 1897.
α particles (Ra.) assuming charge = $+2e$.	By counting α particles and measuring their total charge.	4.65 "	Rutherford & Geiger, <i>P.R.S.</i> , 1908.
Electrolytic ions.	By counting colloid particles.	4.1 "	Perrin, <i>C.R.</i> , 1908.
Charged spray of electrolytic O_2 .	By H. A. Wilson's method, above.	4.7 "	Lattey, <i>P.M.</i> , 1909.
α particles (Polonium); charge = $+2e$.	By counting α particles, and measuring their total charge.	4.79 "	Regener, <i>Berl. Ber.</i> , 1909.
Electrolytic ions.	From Brownian movements.	4.5 "	Brogliè, <i>Le R.</i> ,
Radium rays; negative ions.	By H. A. Wilson's method, above	4.67 "	Begeman, [1909.
		4.77 "	Millikan, <i>P.M.</i> , '17

C.R., *Comptes Rendus*; *Le R.*, *Le Radium*; *P.M.*, *Phil. Mag.*; *P.R.S.*, *Proc. Roy. Soc.*

NUMBER OF MOLECULES IN A GAS

N = the number of molecules in a **gram molecule** of gas (Perrin, *Compt. Rend.*, 1908; Perrin and Dabrowski, *C.R.*, 1909—by observations on colloidal particles). The theoretical value is $N = NE/e = 2.894 \times 10^{14} / (4.77 \times 10^{-10}) = 6.06 \times 10^{23}$.

Method.	Gum mastic.	Gamboge.	Method.	Gum mastic.	Gamboge.
Counting by ultra microscope	$N = 7 \cdot 10^{23}$	$N = 7.05 \cdot 10^{23}$	Brownian movements	$N = 7.3 \cdot 10^{23}$	$N = 7 \cdot 10^{23}$

e/m FOR NEGATIVE ELECTRONS

e/m in E.M.U. gm.⁻¹. Velocities v in cm. sec.⁻¹. For some other values of e/m see J. J. Thomson's "Conduction of Electricity through Gases," and Woltz, *A.d.P.*, 30, 274, 1909. The **mean** of Simon's, Becker's, Classen's, Kaufmann's, Woltz's, Bucherer's, and Bestelmeyer's values is $e/m_0 = 1.772 \times 10^7$ **E.M.U. gm.⁻¹**, where m_0 is the mass of the electron associated with very small velocities. For the variation of e/m with velocity see p. 99. (See also Schuster, *P.R.S.*, 1890.)

e/m	v	Observer.	e/m	v	Observer.
CATHODE RAYS			LENARD RAYS		
1.2×10^7	$2.4 \text{ to } 3.2 \cdot 10^9$	J. J. Thomson, <i>P.M.</i> , 1897	$0.68 \cdot 10^7$	$3.4 \text{ to } 10^7 \cdot 10^9$	Lenard, <i>A.d.P.</i> , 1898
$1.77 \text{ to } 1.8$	—	Kaufmann, <i>A.d.P.</i> , 1897, 1898	INCANDESCENT OXIDES, etc.		
1.86	—	Simon, <i>A.d.P.</i> , 1899	$0.87 \cdot 10^7$	—	J. J. Thomson, <i>P.M.</i> , 1899
1.88	$= e/m_0$		—	—	Owen, <i>P.M.</i> , 1904
1.87	$5.7 \text{ to } 7.5 \cdot 10^9$	Seitz, <i>A.d.P.</i> , 1902	0.56	—	Wehnelt, <i>A.d.P.</i> , 1904
1.84	$3.8 \text{ to } 13$	Starke, <i>V.D.P.G.</i> , 1903	1.5	$0.1 \text{ to } 10 \cdot 10^9$	
1.75	11.1	Becker, <i>A.d.P.</i> , 1905	SECONDARY CORPUSCULAR RAYS, from X-rays incident on platinum		
1.85	$= e/m_0$	Classen, <i>P.Z.</i> , 1908	$1.773 \cdot 10^7$	$= e/m_0$ (on Lorentz's theory)	Bestelmeyer, <i>A.d.P.</i> , 1907.
1.774	$1.9 \cdot 10^9$		ULTRA VIOLET LIGHT ON METAL		
1.767	$3.8 \cdot 10^9$		$0.76 \cdot 10^7$	—	J. J. Thomson, <i>P.M.</i> , 1899
1.771	$= e/m_0$		$1.1 \cdot 10^7$	—	Lenard, <i>A.d.P.</i> , 1900
β RAYS			ZEEMAN EFFECT		
$0.1 \cdot 10^7$	—	Becquerel, <i>Rap. C.P.</i> , 1900	$1.775 \cdot 10^7$	—	Mean of 4 observer's values (see below).
1.77	$= e/m_0$	Kaufmann, <i>Gött. Nachr.</i> , 1901			
1.66	$= e/m_0$ (on Lorentz's theory)	Kaufmann, <i>A.d.P.</i> , 1906			
1.82	$= e/m_0$ (on Abraham's theory)	Kaufmann, <i>A.d.P.</i> , 1906			
1.763	$= e/m_0$	Bucherer, <i>A.d.P.</i> , 1909			
"	$9.5 \text{ to } 20.6 \cdot 10^9$				
1.767	$= e/m_0$	Woltz, <i>A.d.P.</i> , 1909			
"	$15 \text{ to } 21 \cdot 10^9$				

A.d.P., *Ann. der Phys.*; *P.M.*, *Phil. Mag.*; *P.R.S.*, *Proc. Roy. Soc.*; *P.Z.*, *Phys. Zeit.*; *Rap. C.P.*, *Rapports Congrès à Paris*; *V.D.P.G.*, *Verh. Deutschs. Phys. Gesell.*

ELECTRONIC e/m FROM ZEEMAN EFFECT

For a spectrum line of wave-length λ , which becomes a normal triplet with a separation of $\delta\lambda$ in a magnetic field H (in gauss, i.e. E.M.U.), Lorentz has shown that $e/m = 2\pi V \delta\lambda / (\lambda^2 H)$, where V is the velocity of light; e/m is in E.M.U. gm.⁻¹. The values 1.79, 1.77, 1.767, 1.771, mean **1.775 . 10⁷ E.M.U. gm.⁻¹**, agree well with e/m_0 above.

Line.	e/m	Observer.	Line.	e/m	Observer.
Hg 5791, 5770 5461, 4358	$\times 10^7$ 1.72 to 2.80	Blythswood & Marchant, <i>P.M.</i> 1900 [1900	Zn 4810 . . . 4722, 4680. . . He	$\times 10^7$ 2×1.767 1.77	{ Cotton & Weiss, C.R., 1907 Lohmann, <i>P.Z.</i> , 1908
Zn, Cd	1.6	Reese, <i>As. J.</i> , Kent, <i>As. J.</i> , 1901	Hg 5791 . . . " 5770 . . . " 4916 . . .	1.93 2.06 1.81	{ Baeyer & Gehrcke, <i>A.d.P.</i> , 1909
Cd 4678	1.71	Färber, <i>A.d.P.</i> , 1902	" 5790, 5770 " 4916, 4358	1.771	{ Gmelin, <i>A.d.P.</i> , 1909
Zn 4680	1.79	Stettenheimer, <i>A.d.P.</i> , 1907			

ELECTRONIC e/m AND VELOCITY

m_0 is the electromagnetic mass of the negative electron for infinitely small velocities, m the transverse mass for a velocity v ; $v/V = \beta$, where V is the velocity of light. (See Lorentz, *L'Eclairage Électrique*, July, 1905, and "The Theory of Electrons," 1909.) On the theory of Abraham (*Gött. Nachr.*, 1902),

$$\text{transverse mass } m = m_0 3 \left(\frac{1 + \beta^2}{2\beta} \cdot \log \frac{1 + \beta}{1 - \beta} - 1 \right) 4\beta^2$$

β	Infinitely small.	0.1	0.5	0.9	0.99	0.999	0.9999	0.999999
m/m_0 . . .	1.00	1.015	1.12	1.81	3.28	4.96	6.68	10.1

On the theory of Lorentz (*Versl. Kon. Ac. Wet. Am.*, 1904) and the relativity theory of Einstein (*A.d.P.*, 1905), $m = m_0(1 - \beta^2)^{-1/2}$. This theory has been confirmed by the experiments of Bucherer (*A.d.P.*, 1909) and Wolz (*ibid.*), using β rays from Ra with velocities from (9 to 21) $\times 10^9$ cm. per sec. Thus the mass of the negative electron is wholly electromagnetic.

β	m/m_0	β	m/m_0	β	m/m_0	β	m/m_0	β	m/m_0	β	m/m_0	β	m/m_0
0.01	1.005	0.34	1.063	0.48	1.140	0.62	1.274	0.76	1.538	0.90	2.294	0.97	4.113
0.05	1.001	0.36	1.072	0.50	1.155	0.64	1.301	0.78	1.598	0.91	2.412	0.98	5.025
0.10	1.005	0.38	1.081	0.52	1.171	0.66	1.331	0.80	1.667	0.92	2.552	0.99	7.089
0.20	1.020	0.40	1.091	0.54	1.188	0.68	1.364	0.82	1.747	0.93	2.721	0.999	22.36
0.25	1.033	0.42	1.102	0.56	1.207	0.70	1.400	0.84	1.843	0.94	2.931		
0.30	1.048	0.44	1.114	0.58	1.228	0.72	1.441	0.86	1.960	0.95	3.203		
0.32	1.056	0.46	1.126	0.60	1.250	0.74	1.487	0.88	2.105	0.96	3.571		

RH AND v : MAGNETIC DEFLECTION

When negative rays of velocity v are deflected by a uniform magnetic field H (at right angles to their direction) into a circular path of radius R , then $RH = vm/c = v\phi(\beta)/(e/m_0)$, where $\phi(\beta) = (1 - \beta^2)^{-1/2}$ on Lorentz's theory (see above), and $e/m_0 = 1.772 \times 10^7$ E.M.U. gm.⁻¹.

v is in 10^8 cm. sec.⁻¹; RH in gauss cm. **Example.**—If $RH = 1210$ gauss cm.², then $v = 174 \times 10^8$ cm./sec.

RH

v	0	6	12	18	24	30	36	42	48	54	60	66	72	78	84
0	0	33.9	67.8	102	136	170	204	239	274	310	346	382	419	456	494
90	532	572	612	653	695	739	784	830	877	926	977	1030	1090	1150	1210
180	1270	1340	1410	1490	1570	1660	1760	1860	1980	2110	2260	2420	2620	2850	3130
270	3490	3970	4660	5800	8330										

A.d.P., *Ann. der Phys.*; *As. J.*, *Astroph. Journ.*; *C.R.*, *Compt. Rend.*; *P.M.*, *Phil. Mag.*; *P.Z.*, *Phys. Zet.*

α RAYSRANGE AND VELOCITY OF α RAYS

Range in cms. in air at 76 cm. and t° C. (see Bragg and Kleeman, *Phil. Mag.*, 1905). **Initial velocity** (v) in cms./sec. (Rutherford, *Phil. Mag.*, 1906, 1907). Some of the velocities are calculated from the ranges of the α particles; RaC, ThC, and Polonium were observed. **Energy** of RaC α ray = $mv^2/2 = \frac{1}{2}v^2 \cdot 2e \cdot m/e_a = 2 \cdot 06^2 \cdot 10^{18} e / (5 \cdot 07 \cdot 10^3) = 8 \cdot 37 \cdot 10^{14} e = 1 \cdot 3 \cdot 10^{-6}$ ergs = $3 \cdot 1 \cdot 10^{-13}$ calories. Loss of energy in air is proportional to path traversed: thus **initial velocity** of a particle = (velocity of RaC α) $\times \sqrt[3]{r + 1 \cdot 25}$ cm./sec., where r is the range of particle. Also $v = 1 \cdot 077 r^{1/3} \cdot 10^9$ cm./sec. (Geiger, *P.R.S.*, 1910)

α Ray.	Range.	Initial Vel.	Obs.	α Ray.	Range.	Initial Vel.	Obs.
	cms.	cm./sec.			cms.	cm./sec.	
U . . .	634	$1 \cdot 56 \cdot 10^9$	Mc. & R.	Rad.Ac .	48	$1 \cdot 76 \cdot 10^9$	H.
UX . . .	107?	—	Hess.	AcX . . .	655	200 "	H.
Io . . .	28	—	B.	AcEm . .	58	190 "	H.
Ra . . .	$3 \cdot 50/20^\circ\text{C}$.	156 "	B. & K.	AcB . . .	55	186 "	H.
RaEm . .	423	170 "	B. & K.	Th	35	—	—
RaA . . .	483	176 "	B. & K.	Rad.Th .	39	163 "	H.
RaC . . .	706	206 "	B. & K.	ThX . . .	57	189 "	H.
RaF or .	395	—	K.	ThEm . .	55	186 "	H.
Polonium	395	—	K. & M.	ThB . . .	50	179 "	H.
"	386	162 "	L.	ThC . . .	86	225 "	H.

B., Boltwood, *A.J.S.*, May, 1908; B. & K., Bragg & Kleeman, *P.M.*, 1905; H., Hahn, *P.M.*, 1906; Hess, *Wien. Ber.*, 1907; K., Kleeman, *P.M.*, 1906; K. & M., Kucera & Masék, *P.Z.*, 1906; L., Levin, *A.J.S.*, 1906; Mc. & R., McCoy & Ross, *J.A.C.S.*, 1907.

NUMBER OF α PARTICLES FROM Ra

Number of α particles from Ra without its radioactive products = $3 \cdot 4 \cdot 10^{10}$ per gm. per sec. Number of α particles from Ra with its radioactive products = $1 \cdot 36 \cdot 10^{11}$ per gm. per sec. (Rutherford and Geiger, *Proc. Roy. Soc.*, 1908).

 e/m FOR α RAYS

e/m in E.M.U. per gm. $2e/m$ for helium = $2NE/\rho = 4 \cdot 78 \cdot 10^3$ E.M.U./gm. **Mean** for Ra, Pol, RaC = **$4 \cdot 82 \cdot 10^3$ E.M.U. gm $^{-1}$** . Since the α particle is a helium atom with a charge of $2e$, these values should be equal. * Final velocity of rays used.

Subst.	Velocity.*	e/m	Observer.	Subst.	Velocity.*	e/m	Observer.
	cm./sec.	E.M.U.			cm./sec.	E.M.U.	
Ra . . .	$1 \cdot 18$ to $1 \cdot 74 \cdot 10^9$	$4 \cdot 6 \cdot 10^3$	Mackenzie, <i>P.M.</i> , '05	RaA . . .	$1 \cdot 22 \cdot 10^9$	$5 \cdot 6 \cdot 10^3$	Rutherford, <i>P.M.</i> , '06
Pol . . .	$1 \cdot 41 \cdot 10^9$	48 "	Huff (corr d); <i>P.R.S.</i> , '06	AcB . . .	10 "	47 "	
RaC . . .	157 "	$5 \cdot 07$ "	Rutherford, <i>P.M.</i> , '06	ThC . . .	$1 \cdot 98$ "	56 "	Rutherford & Hahn, <i>P.M.</i> , '06

STOPPING POWERS OF MATERIALS

If a layer of air of density ρ and thickness t decreases the range of an α particle by the same amount as aluminium foil of density ρ_a and thickness t_a , then the **atomic stopping power**, S , of Al relative to air is given by $S = 27\rho/14 \cdot 4\rho_a t_a$ = (number of atoms per cm. 2 in air layer)/(number of atoms per cm. 2 in Al foil) (Bragg and Kleeman, *Phil. Mag.*, 1905; Bragg, *Phil. Mag.*, 1906).

Metal.	S.	Metal.	S.	Metal.	S.	Gas.	S.	Gas.	S.
(Air at 20° C., 76 cm.)	100	Ag . . .	317	Ni . . .	246	O ₂ . . .	1055	C ₂ H ₂ . .	111
Al . . .	145	Sn . . .	337	Au . . .	445	N ₂ O . . .	146	Ethylene	135
Cu . . .	243	Pt . . .	416	Pb . . .	427	CO ₂ . . .	147	Benzene	337
		Fe . . .	226	H ₂ . . .	243	CS ₂ . . .	218	Methane	86

A.J.S., *Amer. Journ. Sci.*; *J.A.C.S.*, *Journ. Amer. Chem. Soc.*; *P.M.*, *Phil. Mag.*; *P.R.S.*, *Proc. Roy. Soc.*; *P.Z.*, *Phys. Zeit.*

NUMBER OF IONS MADE BY AN α PARTICLE

Total number of ions produced by the complete absorption of an α particle with various initial velocities. Observer assumed $e = 4.65 \times 10^{-10}$ E.S.U. (Geiger, *Proc. Roy. Soc.*, 1909).

	Ra	RaEm.	RaA	RaC	RaF
Range in air at 20° C., 76 cm. .	3.5 cm.	4.33	4.83	7.06	3.86
Number of ions	1.53×10^5	1.74×10^5	1.87×10^5	2.37×10^5	1.62×10^5

IONS PRODUCED AT DIFFERENT VELOCITIES BY AN α PARTICLE

Number of ions made per mm. of path in air by an α particle from RaC at various distances from its source. Total number = 2.37×10^5 (Geiger, see above).

Distance from RaC in cm.	1	2	3	4	5	6	6.5	7
Ions per mm. of path in air at 12° C. and 76 cm.	2250	2300	2400	2800	3600	5500	7600	4000

TOTAL RELATIVE IONIZATION IN GASES BY α RAYS

I_r = total ionization (relative to air) produced by the complete absorption of α particles in various gases. (B. Bragg, *P.M.*, 1907, used RaC α rays; B. and C., Bragg and Cook, *P.M.*, 1907; L., Laby, *P.R.S.*, 1907, used U α rays; R., Rutherford, *P.M.*, 1899, used U α rays.)

Gas.	I	Gas.	I_r	Gas.	I_r
Air	1.00	Methane	1.16, B. and C.	Et. ether	1.31, B.; 1.29, L.
O ₂	1.09, B.; 1.06, R.	Acetylene	1.26, B.; 1.27, L.	Et. iodide	1.28, B.
N ₂	0.96, B.	Ethylene	1.28, B.	Acetaldehyde	1.05, L.
N ₂ O	1.05, B.; 0.99, L.	Pentane	1.35, B.; 1.345, L.	Chloroform	1.29, B.
NH ₃	1.01, R.; 0.90, L.	Me. alcohol	1.22, B.	Carb. tetra-chloride	1.31, B.
CO ₂	1.08, B.; 1.03, L.	Me. iodide	1.33, B.		
Carbon bi-sulphide	1.37, B.	Et. alcohol	1.23, B.		
		Et. chloride	1.30, B.; 1.18, L.		

RELATIVE VOLUME IONIZATIONS FOR β , γ , AND X RAYS

Relative ionization = $I_r = iP/I_p$, where i is the amount of ionization per unit volume for the gas at a press. p , and I that for air at press. P , the other experimental conditions being the same. In the experiments with γ rays (column headed γ), β rays would also be present. Observers: for β and γ rays, Kleeman, *P.R.S.*, 1907; X rays, C., Crowther, *P.C.P.S.*, 1909; *P.R.S.*, 1909; Mc., McClung, *P.M.*, 1904. I_r for secondary γ rays is much the same as for X rays (see Kleeman, *P.R.S.*, 1909).

Gas.	β	γ	Hard X.	Soft X.	Gas.	β	γ	Hard X.	Soft X.
Air	1.00	1.00	1.00	1.00	Me. alcohol	1.69	1.75	—	—
H ₂	0.16	0.16	0.18, C.	0.01, C.	Me. bromide	3.73	3.81	—	71, C.
O ₂	1.17	1.16	1.17, Mc.	1.3, Mc.	Me. iodide	5.11	5.37	125, C.	145, C.
NH ₃	0.89	0.90	—	—	Chloroform	4.94	4.93	—	—
N ₂ O	1.55	1.55	—	—	CCl ₄	6.28	6.33	71, C.	67, C.
CO ₂	1.60	1.58	1.49, C.	1.57, C.	Et. aldehyde	2.12	2.17	—	—
C ₂ N ₂	1.86	1.71	—	—	Et. bromide	4.41	4.63	118	72, C.
SO ₂	2.25	2.27	4.79, Mc.	11.0, Mc.	Et. chloride	3.24	3.19	17.3, C.	18, C.
CS ₂	3.62	3.66	—	—	Et. ether	4.39	4.29	—	—
Pentane	4.55	4.53	—	—	Et. iodide	5.90	6.17	—	—
Benzene	3.95	3.94	—	—	Ni. carbonyl	—	5.98	97, C.	89, C.
Me. acetate	—	—	3.90, C.	4.95, C.	Hg dimethyl	—	—	—	425, C.

HEAT OF RADIUM

RELATIVE IONIZATION PER UNIT VOLUME BY α RAYS

Relative ionization = (total ionization) \times (stopping power), Metcalfe, *P.M.*, 1909.

Air . 1'00	He . '211	CO . 1'00	HCl . 1'4	Propane 3'05	Pentane 4'83
H ₂ . '233	Br ₂ . 3'9	NO . 1'28	Ethane 2'08	Butane . 4'02	

For calculated **total ionization** when **Röntgen rays** are completely absorbed in various gases, see Crowther, *Proc. Roy. Soc.*, 1909.

HEATING EFFECT OF RADIUM

In calories per sec. per gm. of metallic radium with its radioactive products. E. von Schweidler and Hess, using .795 gm. Ra enclosed in 1 mm. glass + 5 mm. Cu, obtained **.0328 calorie gm⁻¹ sec.⁻¹ = 118 cal. gm.⁻¹ hr.⁻¹** The heating effect of a radioactive substance is proportional to the ionization it produces (Duane, *Le Radium*, 1909). The heat emission continues at temp. of liquid hydrogen (Curie and Dewar, 1903), and is mainly due to the kinetic energy of the α rays (Rutherford, "Radioactivity").

Temp. and press. have no effect on heat emission (Schuster, Eve, and Adams, *Nature*, 1907; Rutherford and Petavel, *B.A. Rep.*, 1907; Schmidt, *P.Z.*, 1908).

Heat.	Observer.	Heat.	Observer.
.0278	Curie and Laborde, <i>C.R.</i> , 1903	25 %	Produced by Ra
.0292	Runge and Precht., <i>Berl. Ber.</i> , 1903	44 %	" " Em + RaB
.0306	Rutherford and Barnes, <i>Nature</i> , 1903; <i>P.M.</i> , 1904	31 %	" " RaC
		.0325	Angström, <i>P.Z.</i> , 1905
		.0372	Precht, <i>A.d.P.</i> , 1906
		.0328	Schweidler and Hess, <i>Wien. Ber.</i> , 1908

HEAT EMISSION FROM RaEm, AND THORIUM

The 6×10^{-4} c.c. of **RaEm** (with its products) in equilibrium with 1 gm. Ra emit .75 of the .0328 calories emitted per sec. by the radium. Thus the total quantity of heat given out by 1 c.c. of RaEm during its whole life = $.75 \times .0328 / (\lambda \times 6 \times 10^{-4}) = 1.9 \times 10^7$ calories.

For old (mineral) **thorium** metal, the heat emitted is 5×10^{-9} calories per sec. per gm. (Pegram and Webb, *Phy. Rev.*, 1908).

RADIUM EMANATION

Γ is the **period of decay** (in days) to half initial activity. Taking $\Gamma = 3.66$ days, then the decay coefficient $\lambda = 2.19 \times 10^{-6}$ sec.⁻¹ (see p. 107).

Γ in days.	Observer, etc.	Γ in days.	Observer, etc.
3.77	Rutherford and Soddy, <i>P.M.</i> , 1903.	3.75	Rümelin, <i>P.M.</i> , 1907.
3.88	Bumstead and Wheeler, <i>A.J.S.</i> , 1904.	3.58	For first 5 days.
3.8 to 4.1	Debierne, <i>C.R.</i> , 1909.	3.75	{ During period 5 to 20 days.
3.86	Sackur, <i>Ber. C.G.</i> , 1905.	3.85	{ 20 to 40 days' old emanation.
		4.4	{ One sample Rutherford and Tuomikoski, <i>P.M.</i> , 1909.

EQUILIBRIUM VOLUME OF RADIUM EMANATION

Final **volume of radium emanation** at 0° C. and 76 cm. Hg in equilibrium with 1 gm. of metallic radium. **Theoretical** volume = (number of radium atoms breaking up per sec.) $\lambda N = 3.4 \times 10^{10} / (2.75 \times 10^{10} \times 2.19 \times 10^{-6}) = 5.64 \times 10^{-4}$ c.c. (Rutherford, "Radioactivity"). The volume of the emanation changes anomalously after it is first formed.

Observed vol.	Observer.	Observed vol.	Observer.
.58 cub. mm.	Rutherford, <i>P.M.</i> , 1908.	.58 cub. mm.	Debierne, <i>C.R.</i> , 1909.
.601 "	Gray & Ramsay, <i>J.C.S.</i> , 1909.		

A.d.P., *Ann. der Phys.*; *A.J.S.*, *Amer. Journ. Sci.*; *B.A. Rep.*, *Brit. Ass. Rep.*; *C.R.*, *Compt. Rend.*; *J.C.S.*, *Journ. Chem. Sci.*; *P.M.*, *Phil. Mag.*; *P.Z.*, *Phys. Zeit.*

VAPOUR PRESSURE OF RADIUM EMANATION

Vapour pressure of liquid RaEm. in cm. Hg; melting-point, -71°C . (R., Rutherford, *Nature*, February, 1909; G. & R., Gray and Ramsay, *J.C.S.*, June, 1909.)

Temp. $^{\circ}\text{C}$	R.	-127°	-101°	-78°	$-65^{\circ} = \text{B.P.}$
Vap. press. cm. Hg		9	5	25	76

Temp. $^{\circ}\text{C}$.	G. & R.	$-70^{\circ}4$	$-62^{\circ} = \text{B.P.}$	$-60^{\circ}6$	$-55^{\circ}8$	$-38^{\circ}5$	$-17^{\circ}7$	$-10^{\circ}2$	$+104^{\circ}5 \text{ crit. t.}$
Vap. press. cm. Hg		50	76	80	100	200	400	500	4745 crit. press.

DIFFUSION OF EMANATIONS

D = coefficient of diffusion (in $\text{cm.}^2 \text{sec.}^{-1}$) of the emanation into the gas stated at the pressure p cm. Hg and temp. $t^{\circ}\text{C}$. indicated. According to J. J. Thomson (*Nature*, November 25, 1909): " D would only vary slowly with atomic weight," and not as the square root of the molecular weight of the emanation, as is assumed in the table below.

Russ finds $pD = \text{const.}$ for AcEm. and for ThEm. Bruhat gives $pD/T^2 = \text{const.}$ for AcEm. between 0° and 20° . (Molec. wgt. ThEm.)/(molec. wgt. AcEm.) = 1.42 (Russ). Mol. wgt. of RaEm. = 222 (Gray & Ramsay, 1910).

Gas.	p. and $t^{\circ}\text{C}$.	D.	Molec. wgt.	Obs.	Gas.	p. and $t^{\circ}\text{C}$.	D.	Molec. wgt.	Obs.
RADIUM EM.					ACTINIUM EM. (contd.)				
Air . . .	76?	'07 to '09	c. 100	R. & B.	Air . . .	—	'112	70	D.
" . . .	76, 10°	'10	—	C. & D.	" . . .	1.4	7.81	—	R.
" . . .	76, 0°	'101	75 to 100	C.	" . . .	76.4	'125	—	"
CO_2 . . .	—	—	180	B. & W.	" . . .	76	'123	—	B.
Diff. of Em. into air compared with $\text{O}_2, \text{CO}_2, \text{SO}_2$ into air . .	—	—	86 to 99	M.	" . . .	$\left\{ \begin{array}{l} 76 \\ 10 \\ 9 \end{array} \right\} 0^{\circ}$	'10	70	"
Em. into H_2 compared with Hg vap. into H_2 . .	250° { 275° {	{ '034 Em. { '037 Hg { '0376 Em. { '0407 Hg	235	P.	THORIUM EM.				
ACTINIUM EM.					Em. into air, compared with $\text{H}_2, \text{O}_2, \text{SO}_2, \text{CO}_2$ into air . .	—	—	c. 90	M.
H_2 . . .	76, 15°	'412	—	B.	Air . . .	—	'09	—	Ruth.
H_2 . . .	76, 10°	'33	—	R.	" . . .	76	'103	—	R.
SO_2 . . .	76, 10°	'062	—	"	" . . .	$\left\{ \begin{array}{l} 8.2 \\ \text{to} \\ 76.1 \end{array} \right\}$	'966 '10 '103	—	"
Argon . . .	to 18°	'106	—	"	" . . .	76	'103	—	"
CO_2 . . .	76, 15°	'073	—	"	Argon . . .	76	'084	—	"
CO_2 . . .	76, 15°	'077	—	B.					

B., Bruhat, *Le Radium*, 1909; B. & W., Bumstead & Wheeler, *A.J.S.*, 1903; C., Chaumont, *Le Radium*, 1909; C. & D., Curie & Danne, *C.R.*, 1903; D., Debiere, *Le Radium*, 1907; M., Makower, *P.M.*, 1905; P., Perkins, *A.J.S.*; R., Russ, *P.M.*, 1909, *Le Radium*, 1909; Ruth., Rutherford, "Radioactivity"; R. & B., Rutherford & Miss Brooks, *C.N.*, 1902.

A.J.S., Amer. Journ. Sci.; *C.N.*, Chem. News; *C.R.*, Compt. Rend.; *J.C.S.*, Journ. Chem. Soc.; *P.M.*, Phil. Mag.

Ra IN ROCKS

EQUILIBRIUM ACTIVITIES IN MINERALS

Relative activity of radioactive products in minerals. Boltwood (*A. J. S.*, April, 1908) found U 2.22 times as active as the Ra alone in minerals (see McCoy and Ross, *A. J. S.*).

Product	U	Io	Ra	RaEm.	RaA	RaB	RaC	RaF	Ac	Total.
Relative activity . .	1	'34	'45	'62	'54	'04?	'91	'46	'28	4.64

3.4×10^{-7} gm. Ra is in equilibrium with 1 gm. U (Rutherford and Boltwood, *A. J. S.*, 1906). 7.3×10^6 gms. U equal in activity 1 gm. of Ra + its products to RaC. *i.e.* Ra just over 30 days old (corrected by Boltwood, *A. J. S.*, 1908).

RADIUM AND THORIUM IN ROCKS

Rutherford and Soddy (*P. M.*, May, 1903) and W. E. Wilson (*Nature*, July, 1903) suggested that the heat liberated by radioactive changes is one of the sources of the Earth's heat. Thus the distribution of radium and thorium in the Earth's crust is of geophysical importance. Loss of heat from the Earth's surface = temperature gradient \times thermal conductivity of crust \times area of Earth's surface = $(1/3200) \times .004 \times 5.1 \times 10^{16} = 6 \times 10^{12}$ calories per sec. Now, elementary radium in radioactive equilibrium (*i.e.* whole U family) gives out 6×10^{-2} cal./sec. gm. (Rutherford §), and therefore 1.1×10^{14} grms. of radium, or $10^{14}/10^{27} = 10^{-13}$ gm. per c.c., throughout the Earth's volume would maintain it at a steady temperature. Thorium contributes 5×10^{-9} cal./sec. gm. The **total heating effect** in calories per gram of rock per hour is for the lava indicated below by *, 30×10^{-10} ; and for the rock indicated by †, 2.9×10^{-10} ; for average igneous rock, 11×10^{-10} .

(See Strutt, *Proc. Roy. Soc.*, 1906-7; Joly, "Radioactivity and Geology," 1909.)

Rock, etc.	Obs.	Ra	Th
		gm. per gm. of rock.	
Igneous rocks	St., 1906	$\times 10^{-12}$	$\times 10^{-5}$
Sedimentary rocks		1.7	—
Sandstone	E. M., 1907	1.1	—
Clays	" "	.16	—
Devonian	" "	.79	—
Ordovician	" "	1 to 4	—
Lavas ejected since 1631 *	" "	.9	—
Lava, Mount Erebus	J., 1909	12.3	2.3
126 igneous rocks	F. F., 1909	2.4	—
64 " "	J., 1909	7.01	—
Italian igneous rocks	other obs.	1.3	1.3
Campbell and Auckland Islands, } N.Z. }	B., 1909	mean	5
St. Gothard Tunnel— granite	F. F., 1909	1.6	igneous
schists and altered sedimentary } rocks }		.5	sedimentary
Simplon Tunnel 	J., 1909	7.7	1.9
Transandine Tunnel †	"	3.4 to 4.9	.5 to 1.2
Calcareous and dolomitic European } rocks }	Fl., 1910	.8	.56
Deep-sea deposits— Globigerina ooze ¹	J., 1910	mean of 7	.16
Radiolarian ooze ²	"	27 samples	<.05
Red clay ³	"	7.2	—
	"	36.7	—
	"	27	—

Extent: —¹ 50, ² 2.5, ³ 51 million square miles. † 1000 feet below the surface. § Assuming that the heat due to each member of the family is proportional to the ionization it produces.

|| Preliminary result. B., Blanc, *P. M.*; E. M., Eve and McIntosh, *P. M.*; F. F., Farr and Florance, *P. M.*; Fl., Fletcher; J., Joly, *P. M.*; S., Strutt (above). *A. J. S.*, *Amer. Journ. Sci.*; *P. M.*, *Phil. Mag.*

RADIUM IN SEA-WATER

In grams per gram of sea-water. Deduced from the observed amount of Ra Em.

Amount.	Place.	Observer.	Amount.	Place.	Observer.
2.3×10^{-15}	—	Strutt, <i>P.R.S.</i> , '06	4×10^{-15}	Nile	Joly, <i>P.M.</i> , 1908
'3-6 "	Mid. N. Atlantic	Eve, <i>P.M.</i> , 1907	14 "	Mediterranean	" " 1909
'9 "	Atlantic	" " 1909	5 "	Indian Ocean	" " "
16 "	"	Joly, <i>P.M.</i> , 1908			

RADIUM EMANATION IN ATMOSPHERE

RaEm. per cubic metre of air, expressed in terms of the number of grams of radium with which it would be in equilibrium. The observers below absorbed the emanation by charcoal.

RaEm.	Place.	Observer.	RaEm.	Place.	Observer.
$24-27 \times 10^{-12}$	Montreal	Eve, <i>P.M.</i> , 1907	$35-350 \times 10^{-12}$	Cam-bridge	Satterly, <i>P.M.</i> , 1908 and 1910
60 "	"	" " 1908	Mean 105 "		
86-200 "	Chicago	Ashman, <i>A.J.S.</i> , '08			

MOBILITIES OF NATURAL IONS IN AIR

Mobility or speed K is in $\text{cm}^2 \text{sec}^{-1} \text{volt}^{-1}$ at room temperature and 76 cm. (see p. 95). The ions are named from their velocities: the small ions are assumed to have the velocity of X-ray ions. (See Pollock, *Science*, 1909; Eve, *Phil. Mag.*, 19, 1910; Lusby, *Proc. Camb. Phil. Soc.*, 1910.)

Ion.	Mean K .	Observer.	Ion.	Mean K .	Observer.
Small . . .	$\begin{cases} K_+ = 1.4 \\ K_- = 1.7 \end{cases}$	Langevin, '03	Large . .	'0003	Langevin, <i>C.R.</i> , '05
Intermediate	0.01	Mean	Large . .	'0003 *	Pollock, 1908
			Large . .	'0008 †	" "

* Humidity, 19 grms. H_2O per cubic metre. † 5 grm. H_2O per cubic metre of air. Pollock, *Austl. Ass. Adv. Sci.*, 1908.

ELECTRIC ARCS

Mrs. Ayrton's formula for carbon arcs, $E = a + \beta l + \frac{\gamma + \delta l}{i}$, has been shown by

Guye and Zébrakoff (*Compt. Rend.*, 1907) to hold for short stable arcs between metals. E is the voltage across the arc, i is the current in amperes, and l the length in mms. of the arc in air at atmospheric pressure. Mrs. Ayrton's formula does not hold for very long arcs, nor for cored carbons. For stability, an arc requires an external resistance R which must be less than $\frac{\{E_x - (a + \beta l)\}^2}{4(\gamma + \delta l)}$ ohms, where E_x is the total available voltage; or E_x must exceed $a + \beta l + 2\sqrt{R(\gamma + \delta l)}$. If R is too small the arc hisses, in which case the current is independent of the voltage across the terminals. The constants for carbon refer only to the particular sizes and quality used by Mrs. Ayrton.

(See J. J. Thomson, "Conduction of Electricity through Gases.")

Metal.	a	β	γ	δ	Metal.	a	β	γ	δ
C . . .	38.88	2.074	11.66	10.54	Pd . . .	21.64	3.70	0	21.78
Fe . . .	15.73	2.52	9.44	15.02	Ag . . .	14.19	3.64	11.36	19.01
Ni . . .	17.14	3.89	0	17.48	Pt . . .	24.29	4.80	0	20.23
Co . . .	20.71	2.05	2.07	10.12	Au . . .	20.82	4.62	12.17	20.97
Cu . . .	21.38	3.03	10.69	15.24					

A.J.S., Amer. Journ. Sci.; *C.R.*, Compt. Rend.; *P.M.*, Phil. Mag.; *P.R.S.*, Proc. Roy. Soc.

ATOMIC CONSTANTS

ATOMIC AND RADIOACTIVITY CONSTANTS

References : J. J. Thomson's "Conduction of Electricity through Gases," Rutherford's "Radioactivity," H. A. Lorentz, *Éclairage Electrique*, 44, 1905, "Theory of Electrons," 1909, Jeans' "Dynamical Theory of Gases," and Millikan, *P.M.*, 1917.

Symbol.	Definition.	Value.
e.	Ionic charge , half charge on an α particle	$4.77 \cdot 10^{-10}$ E.S.U.; $1.59 \cdot 10^{-20}$
NE	Total charge carried in electrolysis by the atoms in $\frac{1}{2}$ c.c. of gas— For ideal gas at 0° and 76 cm.	[E. M. U.; $1.59 \cdot 10^{-19}$ coulombs]
	„ oxygen „ „ „	$1.2913 \cdot 10^{10}$ E.S.U. cm. $^{-3}$; 4304 E.M.U. cm. $^{-3}$
	„ hydrogen „ „ „	$1.292 \cdot 10^{10}$ E.S.U. cm. $^{-3}$; 4308 E.M.U. cm. $^{-3}$
N_mE	Total charge carried by $\frac{1}{2}$ (gm. molecule) of hydrogen ions	$1.290 \cdot 10^{10}$ E.S.U. cm. $^{-3}$; 4300 E.M.U. cm. $^{-3}$
N	Number of molecules per c.c. of a gas at 0° C. and 76 cm. = $NE/e = 1.29 \cdot 10^{20}/4.77$	$2.894 \cdot 10^{14}$ E.S.U. cm. $^{-3}$; $9.647 \cdot 10^3$ E.M.U. cm. $^{-3}$
N_m	Number of molecules in 1 gm. molecule of gas	$2.705 \cdot 10^{19}$ cm. $^{-3}$
e/m₀	Ratio of charge to electromagnetic mass for the negative electron at small velocities	$6.062 \cdot 10^{23}$ gm. $^{-1}$ $5.31 \cdot 10^{17}$ E.S.U. gm. $^{-1}$; $1.77 \cdot 10^7$ E.M.U. gm. $^{-1}$
E/m_H	The same ratio for the hydrogen ion in electrolysis = $107.88/(\cdot 00111827 \times 1.008)$	9.571 E.M.U. gm. $^{-1}$; $95,706$ coulombs gm. $^{-1}$
e/m_{α}	The same ratio for the α particle	$4.8 \cdot 10^3$ E.M.U. gm. $^{-1}$
2e/m_{He}	Calculated for helium = $2NE/\rho = 2 \times .43 \cdot 10^{-6}/(2 \times 8.987)$	$4.78 \cdot 10^3$ E.M.U. gm. $^{-1}$
m₀	Electromagnetic mass of negative electron for small velocities = $e/(e/m_0)$	$8.8 \cdot 10^{-28}$ gm.
m_H	Mass of hydrogen atom = $\rho/2N$	$1.66 \cdot 10^{-24}$ gm.
m_{α}	Mass of a particle , i.e. of helium atom	$6.56 \cdot 10^{-24}$ gm.
m_H/m₀	Number of electrons equal in mass to hydrogen atom = $(m_{He})/(m_0E)$	1850
$\alpha\theta$	Energy of a gas molecule at θ° C. = $3p/2N$	$\alpha = 2.02 \cdot 10^{-18}$ ergs/degree
R	For 1 gm. of oxygen, $R = pv/\theta = 1.0132 \cdot 10^6/(273.09 \cdot 1.429 \cdot 10^{-3})$. Press. in dynes/cm. 2 ; volume in c.c. (see p. 5) For 1 gm. molecule of an ideal gas, $R = 22.412/273.09$. Press. in atmos. = 76 cm. Hg ($g = 980.6$); vol. in litres (D. Berthelot, <i>Trav. et Mém. Bur. Intl.</i>)	$\{2.5963 \cdot 10^6$ cm. 2 /sec. 2 $\{2.5963 \cdot 10^6$ ergs/gm. mol. $\cdot 08207$ litre atm./gm. mol.
a.	The radius of a negative electron = $2/3 \cdot e \cdot e/m_0$	$1.85 \cdot 10^{-13}$ cm.
	The diameter of a hydrogen molecule (Sutherland (after Jeans), <i>Phil. Mag.</i> , 1910)	$2.17 \cdot 10^{-8}$ cm. (see p. 33)
Heat given out by 1 gm. of metallic radium with its products:		$\cdot 0328$ cal./sec.; 118 cal./hr.
Number of α particles emitted by 1 gm. radium without products		$3.4 \cdot 10^{10}$ gm. $^{-1}$ sec. $^{-1}$
Initial velocity of a particle from RaC		$2.06 \cdot 10^9$ cm./sec.
Initial energy of a particle from RaC = $mv^2/2 = v^2e/(2e/m_0) = 2.06^2 \cdot 10^{18} \times 1.57 \cdot 10^{-20}/(2 \times 5.07 \cdot 10^3)$		$1.3 \cdot 10^{-5}$ ergs; $3.1 \cdot 10^{-13}$ cal.
Total number of ions produced in air by an α ray (RaC)		$2.37 \cdot 10^5$
Volume of helium at 0° and 76 cm. produced by 1 gm. radium		$5.17 \cdot 10^{-9}$ c.c. / (sec. gm.), or 163 mm. 3 /(yr. gm.)
Calculated volume = $4 \times$ number of α rays emitted/N = $4 \cdot 3.4 \cdot 10^{-9}/2.75$		$4.94 \cdot 10^{-9}$ c.c. / (sec. gm.); 156 mm. 3 / (yr. gm.)
Number of β particles emitted per sec. by the RaC in equilibrium with 1 gm. Ra (Makower, <i>Phil. Mag.</i> , 1909)		$5 \cdot 10^{10}$ gm. $^{-1}$ sec. $^{-1}$

CONSTANTS OF RADIOACTIVE SUBSTANCES

Atomic weights: O = 16, U = 238.2, Ra = 226.0, Th = 232.4.

Rate of decay: If I is the radioactivity of a substance at a time t , then $I = I_0 e^{-\lambda t}$, where I_0 is the initial activity when $t = 0$. λ is given below in sec.^{-1} . If Γ is the period in which the activity decreases to half its initial value (i.e. $I/I_0 = \frac{1}{2}$), then $\lambda = .69315/\Gamma \text{ sec.}^{-1}$. Γ is given below in secs. (s.), mins. (m.), hrs. (h.), days (d.), or years (y.).

Coefficients of absorption Λ are given in cm.^{-1} for β rays in Al foil and for γ rays in lead foil. If J_0 is the intensity of the rays incident on foil of thickness $d \text{ cm.}$, and J is the intensity of the emergent rays, then $J = J_0 e^{-\Lambda d}$.

(See Rutherford's "Radioactive Substances," Camb. Univ. Press, and Wendt, *Phys. Rev.*, 1916, for a complete table.)

Substance.	λ in sec.^{-1} .	Half-period. Γ	Rays emitted.	Absorptn. Coef. in cm.^{-1} .	
				β Rays. Λ_{Al}	γ Rays. Λ_{Pb}
U	$4.3 \cdot 10^{-18}$	$6 \cdot 10^9 \text{ y.}$	α	—	—
Rad. U	—	sevl. y.	—	—	—
U.X.	$3.7 \cdot 10^{-7}$	21.5 d.	β, γ	14.4 and 510	.72
Io	$2 \cdot 10^{-12}$	$c. 10^4 \text{ y.}$	α	—	—
Ra	$1.1 \cdot 10^{-11}$	2000 y.	α, β	312	—
RaEm	$2.08 \cdot 10^{-6}$	3.85 d.	α	—	—
RaA.	$3.85 \cdot 10^{-3}$	3 m.	α	—	—
RaB.	$4.33 \cdot 10^{-4}$	26.7 m.	β	13 to 890	—
RaC ₁	$5.93 \cdot 10^{-4}$	19.5 m.	α	13 to 53	.46 to .57
RaC ₂	—	1.38 m.	β, γ	—	—
Rad.	$1.8 \cdot 10^{-9}$	17 y.	β	—	—
RaE ₁	$1.3 \cdot 10^{-6}$	6.2 d.	β	—	—
RaE ₂	$1.7 \cdot 10^{-6}$	4.8 d.	β	44	—
RaF (Polonium)	$5.73 \cdot 10^{-8}$	140 d.	α	—	—
Ac	—	30 y.	rayless	—	—
Rad. Ac	$4.1 \cdot 10^{-7}$	19.5 d.	α, β	170	—
AcX.	$7.6 \cdot 10^{-7}$	10–11 d.	α	—	—
AcEm	$1.8 \cdot 10^{-1}$	3.9 s.	α	—	—
AcA.	—	0.002 s.	α	—	—
AcB.	$3.20 \cdot 10^{-4}$	36.1 m.	β	—	—
AcC.	$5.37 \cdot 10^{-3}$	2.15 m.	α	—	—
AcD.	$2.26 \cdot 10^{-3}$	4.71 m.	β	29	2.0 to 3.6
Th	$7 \cdot 10^{-19}$	$3 \cdot 10^{10} \text{ y.}$	α	—	—
MesoTh 1	$4.0 \cdot 10^{-9}$	5.5 y.	rayless	—	—
MesoTh 2	$3.1 \cdot 10^{-3}$	6.2 h.	β, γ	20.2 to 38.5	.5
Rad. Th	$1.09 \cdot 10^{-8}$	737 d.	α	—	—
ThX.	$2.17 \cdot 10^{-6}$	3.71 d.	α	—	—
ThEm	$1.31 \cdot 10^{-2}$	53 s.	α	—	—
ThA.	—	—	α	140	—
ThB.	$1.81 \cdot 10^{-5}$	10.6 h.	β	—	—
ThC ₁	$2.10 \cdot 10^{-4}$	55 m.	α	—	—
ThC ₂	—	some secs.	α	—	—
ThD.	$3.7 \cdot 10^{-3}$	3.1 m.	β	15.7	.46 to .57

PROPERTIES OF RADIOACTIVE SUBSTANCES

Substance.	Properties.	Substance.	Properties.
U . . .	Sol. in excess of am. carb. Nitrate soluble in ether and acetone.		Carried down by PbCO_3 , and by SnCl_2 with Hg and Te. RaD , E_1 , F_2 , and F can be separated by electrolysis.
Rad.U . .	Carried down by BaSO_4 and ferric hydrate. Soluble in HCl.		
U.X . . .	Less volatile than U. Volatile in electric arc. Insoluble in excess of am. carb. Soluble in water and ether. Carried down by barium sulphate, by moist ferric hydrate, and by animal charcoal.	Ac . . .	Produces helium. Precipitated by oxalic acid in acid solutions. Oxalate insoluble in HF; accompanies thorium and rare earths.
		Rad.Ac . .	Slightly volatile at high temps. Insoluble in NH_4OH . Separated from Ac by electrolysis, by fractional precipitation, by ammonia, and by animal charcoal.
Io . . .	Soluble in excess of am. oxalate. Carried down by H_2O_2 in presence of U salts.	AcX . . .	Deposited by electrolysis in alkaline solution. Not precipitated by NH_4OH .
Ra . . .	Characteristic spectrum. Spontaneously luminous. Analogous to Ba. RaCl_2 and RaBr_2 are less soluble than BaCl_2 and BaBr_2 .	AcEm . . .	Behaves as inert gas. Coef. of diffusion in air 0.11. Condenses at -120°C .
RaEm . . .	One of group of inert gases. Characteristic spectrum. Coef. of diffusion in air = 0.1 (see p. 103). Mol. wt. = 218.	AcA . . .	Volatile below 400°C . Soluble in NH_4OH and strong acids.
RaA . . .	Behaves as a solid. Deposited on cathode in an electric field. Volatile at 800 – 900°C . Soluble in strong acids.	AcB . . .	Volatile below 700°C . Soluble in NH_4OH and strong acids. Deposited by electrolysis of active deposit on the cathode in HCl.
RaB . . .	Like RaA. Volatile at 600 – 700°C . Precipitated by BaSO_4 .	Th . . .	Volatile in electric arc. Colourless salts not spontaneously phosphorescent. Salts pptd. by NH_4OH and oxalic acid.
RaC . . .	Physically like RaA. Volatile at 800 – 1300°C . Chemically, like RaB. Deposited on Cu and Ni. Carried down with precipitated copper. Perhaps a mixture of 2 or 3 products.	Rad.Th . .	Carried down by hydrates, precipitated by NH_4OH .
RaD . . .	Volatile below 1000°C . Soluble in strong acids. Reactions analogous to those of Pb.	ThX . . .	Soluble in NH_4OH . Carried down by iron. Deposited by electrolysis in alkaline soln.
RaE₁ . . .	Volatile at red heat. Soluble in cold acetic acid. Reactions analogous to those of Pb.	ThEm . . .	Inert gas. Condenses just above -120°C . Coefficient of diffusion in air = .10.
RaE₂ . . .	Not volatile at red heat. Reactions analogous to those of bismuth.	ThA . . .	Volatile under 630°C . Soluble in strong acids.
RaF(Pol.)	Volatile towards 1000°C . Deposited from its solutions on Bi, Cu, Sb, Ag, Pt.	ThB . . .	Volatile below 730°C . Like ThA. Deposited on Ni. Separated from ThA by electrolysis.
		ThC . . .	Like ThB.

PHYSICAL CONSTANTS OF CHEMICAL COMPOUNDS

For properties of the **elements**, see: density, p. 20; melting and boiling points, p. 48; solubility in water, p. 124. **Metallo-organic** compounds are given under "Organic Compounds," p. 118.

Formulae.—Hydrated forms (which are often crystalline) are indicated thus: $\text{CaI}_2(\text{and} + 6\text{H}_2\text{O})$; the properties given are for the anhydrous substance.

Formula (Molecular) Weights are calculated with atomic weights for 1911 (p. 1).

Densities.—When no temp is given, grams. per c.c. at 15° may be assumed. When preceded by "A" the density is relative to that of air ('001293 gram per c.c. at 0° and 760 mms.). To convert this into a density relative to $\text{O} = 16$, multiply by 14.47. For those gaseous densities known with accuracy, see p. 26. Other densities on pp. 20-26.

Melting and Boiling Points are for anhydrous substances at 760 mms. mercury unless some other conditions are specified. T = temp. of transition or pseudo-"melting" point of hydrated substance. For fats and waxes, see p. 50.

Solubilities are given as grams of substance in 100 grams of water at the temp. stated. "g" indicates grams per 100 grams of solution. "V" means volumes of substance at 0° and 760 mms. per 100 volumes of water at the temp. stated. "Soluble" infers solubility in either hot or cold water; "insoluble" indicates solubility in neither. (See also pp. 124, 125.)

For more complete tables, see Van Nostrand's "Chemical Annual" and Biedermann's "Chemiker-Kalender" for current year; Dammer's "Handbuch der Anorganischen Chemie"; Beilstein's "Handbuch der Organischen Chemie"; Watts' "Dictionary of Chemistry"; and F. W. Clarke's "Specific Gravities."

INORGANIC COMPOUNDS

Formula, formula (molecular) weight, density, melting and boiling points, and solubility in water.

Substance and Formula.	Formula weight ($\text{O} = 16$).	Density, gms./c.c.	Melting Point, $^\circ\text{C}$.	Boiling Point, $^\circ\text{C}$.	Solubility in Water.
Aluminium —		at./temp.	at./mms.	at./mms.	at./temp.
bromide, $\text{Al}_2\text{Br}_6(\text{and} + 12\text{H}_2\text{O})$	533.7	$\left\{ \begin{array}{l} 2.54; \\ \text{A. } 18.62 \end{array} \right\}$	93°	$263^\circ/747$	soluble
chloride, $\text{Al}_2\text{Cl}_6(\text{and} + 12\text{H}_2\text{O})$	267.0	A. $9.34/400^\circ$	$190^\circ/1910$	$182^\circ/752$	$41/15^\circ(\rho)$
iodide, $\text{Al}_2\text{I}_6(\text{and} + 12\text{H}_2\text{O})$	815.7	$\left\{ \begin{array}{l} 2.63; \\ \text{A. } 27 \end{array} \right\}$	185°	360°	soluble
nitrate, $\text{Al}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$	375.3	—	T = 73°	dec. 134°	v. soluble
oxide, Al_2O_3	102.2	3.7 - 4	2200	—	insoluble
phosphate, AlPO_4	122.1	2.59	infusible	—	insoluble
sulphate, $\text{Al}_2(\text{SO}_4)_3 \cdot 18\text{H}_2\text{O}$	666.7	1.62	decomp.	—	$36/20^\circ$
Potassium alum, $\text{Al}_2(\text{SO}_4)_3 \cdot \text{K}_2\text{SO}_4 \cdot 24\text{H}_2\text{O}$	949.1	$1.757/20^\circ$	$84^\circ.5$	$\left\{ \begin{array}{l} 23\text{H}_2\text{O} \\ \text{at } 190^\circ \end{array} \right\}$	$\left\{ \begin{array}{l} 9.6/15^\circ \\ 357/100^\circ \end{array} \right\}$
Ammonium —					
ammonia, NH_3	17.03	$\left\{ \begin{array}{l} (\text{liq.}) .623/0^\circ \\ \text{A. } .5896 \end{array} \right\}$	- 75	- 33.5	see p. 124.
acetate, $\text{NH}_4\text{C}_2\text{H}_3\text{O}_2$	77.07	—	89	—	$148/4^\circ$
arsenate, $(\text{NH}_4)_3\text{AsO}_4 \cdot 3\text{H}_2\text{O}$	247.1	—	—	—	soluble
bromide, NH_4Br	97.96	$\left\{ \begin{array}{l} 2.33/15^\circ \\ \text{A. } 1.64/440^\circ \end{array} \right\}$	diss.	—	$\left\{ \begin{array}{l} 66/10^\circ \\ 128/100^\circ \\ 100/15^\circ \end{array} \right\}$
carbonate, $(\text{NH}_4)_2\text{CO}_3 \cdot \text{H}_2\text{O}$	114.1	—	diss. 85°	—	$100/15^\circ$
chloride, NH_4Cl	53.50	$\left\{ \begin{array}{l} 1.52/17^\circ \\ \text{A. } .89 \end{array} \right\}$	diss. 350°	—	$\left\{ \begin{array}{l} 35/15^\circ; \\ \text{see p. 125.} \end{array} \right\}$
chloroplatinate, $(\text{NH}_4)_2\text{PtCl}_6$	444.0	3.06	decomp.	—	$67/20^\circ$
chromate, $(\text{NH}_4)_2\text{CrO}_4$	152.2	$1.88/11^\circ$	decomp.	—	decomp.
iodide, NH_4I	145.0	2.5	sublimes	—	v. soluble
molybdate, $(\text{NH}_4)_2\text{MoO}_4$	196.1	$2.4 - 2.9$	decomp.	—	decomp.
nitrate, NH_4NO_3	80.05	$1.72/15^\circ$	152°	dec. 210°	$200/18^\circ$

dec. or decomp. = decomposes; diss. = dissociates; v. = very; wh. = white.

PHYSICAL CONSTANTS

INORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, ° C.	Boiling Point, ° C.	Solubility in Water.
Ammonium (<i>contd.</i>) —					
nitrite, NH_4NO_2	64.05	at./temp. 1.7	at./mms. decomp.	at./mms. —	at./temp. soluble
oxalate, $(\text{NH}_4)_2\text{C}_2\text{O}_4 \cdot \text{H}_2\text{O}$	142.1	1.5	—	—	4/15°
persulphate, $(\text{NH}_4)_2\text{S}_2\text{O}_8$	228.2	—	decomp.	—	58/6°
phosphomolybdate, $(\text{NH}_4)_3\text{PO}_4 \cdot 12\text{MoO}_3 \cdot 3\text{H}_2\text{O}$	1931	—	—	—	.03/15°
sulphate, $(\text{NH}_4)_2\text{SO}_4$	132.2	1.77/20°	140°	dec. 280°	76/20°
sulphocyanate, NH_4CNS	76.12	1.31/13°	159	dec. 170°	162/20°
Antimony —					
bromide, SbBr_3	360.0	4.15/23°	93	280°	decomp.
chloride, tri-, SbCl_3	226.6	{ 3.06/26° A. 8.1 }	73.2	223	{ 816/15° ∞/72°
„ penta-, SbCl_5	297.5	2.35/20°	— 6	102°/68	decomp.
hydride, SbH_3	123.2	A. 4.3/15°	— 91.5	— 18	20 V.
iodide, tri-, SbI_3	501.0	{ 4.85/26° A. 17.6 }	167 subl. 114°	401	decomp.
oxide, tri-, Sb_2O_3	288.4	5.2-5.7	red heat	1550	.002/15°
„ tetr-, Sb_2O_4	304.4	4.07	O/800°	—	insoluble
„ pent-, Sb_2O_5	320.4	3.8	O/300°	O ₂ /800°	insoluble
potassium tartrate, $\text{K}(\text{SbO})\text{C}_4\text{H}_4\text{O}_6 \cdot \frac{1}{2}\text{H}_2\text{O}$	332.3	2.6	$\frac{1}{2}\text{H}_2\text{O}$ /100°	decomp.	{ 5/9° 36/100°
sulphide, tri-, Sb_2S_3	336.6	4.65	fusible	volatilizes	insoluble
„ penta-, Sb_2S_5	400.7	4.12/0°	fusible	—	insoluble
Arsenic —					
bromide, AsBr_3	314.7	{ 3.7/15° A. 10.91 }	31°	221°	decomp.
chloride, AsCl_3	181.3	2.2/0°; A. 6.3	— 18	130.2	decomp.
fluoride, tri-, AsF_3	132.0	2.7; A. 4.57	— 8.5	63	decomp.
„ penta-, AsF_5	170.0	A. 4.15	— 80	— 53	soluble
hydride, AsH_3	77.98	A. 2.7	— 113	— 54.8	slghtly sol.
iodide, di-, AsI_2	328.8	—	—	—	—
„ tri-, AsI_3	455.7	4.4/13°	146	{ 394-414 V.D. 16.1 }	30/100°
„ pent-, AsI_5	709.6	3.93	70	—	decomp.
oxide, tri-, As_2O_3	197.9	3.6-4.1	subl. 218°	V.D. 13.8	17/16°
„ pent-, As_2O_5	229.9	3.9-4.2	red heat	decomp.	245/12°
Barium —					
bromide, $\text{BaBr}_2 \cdot 2\text{H}_2\text{O}$	333.2	3.85/24°	anhy. 880°	2H ₂ O/100°	103/15°
carbonate, BaCO_3	197.4	4.3	795°	dec. 1450°	.002/18°
chloride, $\text{BaCl}_2 \cdot 2\text{H}_2\text{O}$	244.3	3.1/24°	anhy. 960°	2H ₂ O/113°	see p. 125.
hydride, BaH_2	139.4	4.2/0°	volatile	1400°	decomp.
iodide, BaI_2	391.2	4.92	740°	—	170/0°
nitrate, $\text{Ba}(\text{NO}_3)_2$	261.4	3.24/23°	575	—	5/0°
oxide, BaO	153.4	4.7-5.5	BaO ₂ /450°	—	1.5/0°
„ per-, BaO_2	169.4	4.96	BaO/450°	—	insoluble
sulphate, BaSO_4	233.4	c. 4.5	infusible	—	.03/23/18°
Beryllium —					
bromide, BeBr_2	168.9	—	601°	—	soluble
chloride, BeCl_2	80.02	—	c. 600	—	v. soluble
sulphate, $\text{BeSO}_4 \cdot 4\text{H}_2\text{O}$	177.2	1.7/10°	dec. r. ht.	2H ₂ O/100°	44/30°

anhy. = anhydrous; dec. or decomp. = decomposes; r. ht. = red heat; subl. = sublimes;
v. = very; V.D. = vapour density; ∞ = soluble in all proportions.

INORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.	Solubility in Water.
Bismuth—		at./temp.	at./mms.	at./mms.	at./temp.
bromide, BiBr ₃	447.76	5.6	200°–215°	453°	decomp.
chloride, tri-, BiCl ₃	314.38	{ 4.6/11° A. 11.35 }	227	429	decomp.
nitrate, Bi(NO ₃) ₃ ·5H ₂ O	484.11	2.8	74	5H ₂ O/80°	decomp.
oxide, Bi ₂ O ₃	464.0	8.8 – 9	820–860	—	insoluble
sulphide, Bi ₂ S ₃	512.21	7 – 7.8	decomp.	—	insoluble
Boron—					
chloride, BCl ₃	117.38	1.35/0°; A. 4/17°	—	18°–2	decomp.
fluoride, BF ₃	68.0	A. 2.3	–127°	–101	decomp.
oxide, B ₂ O ₃	70.0	1.83/4°	577	—	16/102°
Borax. <i>See</i> Sodium borate.					
Boric acid, H ₃ BO ₃	62.0	1.43/15°	184–186	H ₂ O/100°	4/18°
Cadmium—					
bromide, CdBr ₂	272.24	4.7–4.9/14°	571	806–812	48.9/18° <i>p</i> .
chloride, CdCl ₂	183.32	3.6/15°	590	c. 900	140/20°
nitrate, Cd(NO ₃) ₂ ·4H ₂ O	308.48	2.4	59.5	132	127/18°
oxide, CdO	128.4	6.9–8.1	infusible	—	insoluble
sulphate, anhy. CdSO ₄	208.47	4.7/15°	1000°	—	59/23°
„ hydr. 3CdSO ₄ ·8H ₂ O	769.54	3.05	—	—	see p. 125.
Cæsium—					
carbonate, Cs ₂ CO ₃	325.62	—	<red heat	dec. 610°	v. soluble
chloride, CsCl	168.27	3.97/20°	631°	sublimes	174/10°
hydride, CsH	133.82	2.7	decomp.	—	decomp.
hydroxide, CsOH	149.82	4.02	red heat	—	soluble
nitrate, CsNO ₃	194.82	3.69/28°	414°	decomp.	15/10°
Calcium—					
bromide, CaBr ₂	199.93	3.3/20°	760	c. 800°	125/0°
carbonate, CaCO ₃	100.09	2.7–2.9	dec. 825°	—	0.018 cold
chloride, anhy. CaCl ₂	111.0	2.3/20°	780°	4H ₂ O/30°	63/10°
„ hydr. CaCl ₂ ·6H ₂ O	219.1	1.65	29	6H ₂ O/200°	96/0°
hydride, CaH ₂	42.11	1.7	—	—	decomp.
hydroxide, Ca(OH) ₂	74.11	2.08	—	—	see p. 125.
iodide, CaI ₂ (and +6H ₂ O)	293.1	4.9/20°	740	c. 710	192/0°
nitrate, Ca(NO ₃) ₂ ·4H ₂ O	236.17	1.82	561	dec. 132°	54.8/18°
oxide, CaO	56.09	3.08	abt. 2000	—	13/0°
phosphate, Ca ₃ (PO ₄) ₂	310.3	3.2	—	—	0.03–0.08
sulphate, CaSO ₄	136.16	2.96	—	—	18/0°
Carbon—					
chloride, tetra-, CCl ₄	153.84	1.582/21°	–23°–8	76°–7	insoluble
oxide, sub- (1906), C ₃ O ₂	68.00	—	—	7°/761	*
„ mon-, CO	28.00	A. .967	–207	–190	see p. 124.
„ di-, CO ₂	44.00	liq. .772/20°†	–65	–78.2	see p. 124.
sulphide, mono- CS	44.07	1.6–1.83	—	—	—
„ bi-, CS ₂	76.14	1.292/0°	–110	46.2	2/0°
Cerium—					
chloride (cerous), CeCl ₃	246.63	3.88/15°–5	v. fusible	—	soluble
oxide (cerous), Ce ₂ O ₃	328.5	6.9–7	—	—	insoluble
„ (ceric), CeO ₂	172.25	6.74	—	—	insoluble
sulphate (cerous), Ce ₂ (SO ₄) ₃ ·8H ₂ O	712.84	3.22	8H ₂ O/630°	—	16.5/0°
Chlorine—					
oxide, mon-, Cl ₂ O	86.92	{ liq. 3.87 A. 3.007 }	explosive	–19	200V/0°

* Forms malonic acid.

† Behn, *Ann. d. Phys.*, 1900.

anhy. = anhydrous ;

dec. or decomp. = decomposes ; hydr. = hydrated ; liq. = liquid ; v. = very.

PHYSICAL CONSTANTS

INORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.	Solubility in Water.
Chlorine (<i>contd.</i>)—		at./temp.	at./mins.	at./mms.	at./temp.
oxide, di-, ClO_2	67.46	1.5; A. 2.3	-76°	9.9°/731	20V/4°
Chromium —					
chloride (chromous), CrCl_2	122.92	2.75/14°	—	—	v. soluble
„ (chromic), CrCl_3	158.38	{ 2.76/15° A. 11/1200° }	—	c. 1300°	slightly sol.
oxide, Cr_2O_3	152.0	5.04	white heat	—	insoluble
„ tri-, CrO_3	100.0	2.74	190	decomp.	62.1/0°(p)
sulphate, $\text{Cr}_2(\text{SO}_4)_3 \cdot 15\text{H}_2\text{O}$	662.65	1.867/17°	15H ₂ O/100°	—	120/20°
Cobalt —					
cobaltous chloride,					
$\text{CoCl}_2(\text{and} + 6\text{H}_2\text{O})$	129.9	2.94	subl. c. 87°	—	29.5/0°
„ hydrate, $\text{Co}(\text{OH})_2$	93.02	3.6/15°	—	—	insoluble
„ oxide, CoO	74.98	5.7	dec. 100°	—	insoluble
„ sulphate,					
$\text{CoSO}_4 \cdot 7\text{H}_2\text{O}$	281.2	1.918/15°	96°-8	—	26/3°
cobaltic chloride, CoCl_3	165.35	2.94	sublimes	—	soluble
„ oxide, Co_2O_3	165.95	5.1	dec. r. ht.	—	insoluble
„ sulphate, $\text{Co}_2(\text{SO}_4)_3$	406.15	—	—	—	soluble
Columbium. See Niobium.					
Copper —					
cuprous chloride, Cu_2Cl_2	198.06	{ 3.7 A. 6.6/1690° }	410	c. 1000°	insoluble
„ oxide, Cu_2O	143.14	5.8-6.1	red heat	—	insoluble
cupric chloride, CuCl_2	134.49	3.05	498	decomp.	75/17°
„ nitrate, $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$	241.64	2.17	114.5	{ 170° dec. r. ht. }	{ 60/25°(p) insoluble }
„ oxide, CuO	79.57	6.30	—	—	insoluble
„ sulphate, $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$	249.65	2.28/15°	{ 4H ₂ O/100° 5H ₂ O/240° }	dec. r. ht.	see p. 125.
Cyanogen, C_2N_2	52.02	{ liq. .866/17° A. 1.806 }	-35°	-207°	4.5 V/20°
Erbium —					
oxide, Er_2O_3	382.8	8.6	infusible	—	insoluble
sulphate, $\text{Er}_2(\text{SO}_4)_3 \cdot 8\text{H}_2\text{O}$	767.14	3.18	dec. 950°	—	23/20°
Gadolinium —					
sulphate, $\text{Gd}_2(\text{SO}_4)_3$	602.81	4.14/15°	—	—	2.3/34°
Gallium —					
chloride, tri-, GaCl_3	176.28	A. 12.2/240°	75°-5	220	decomp.
Germanium —					
chloride, tetra-, GeCl_4	214.34	1.89/18°	—	86	decomp.
oxide, di-, GeO_2	104.5	4.70/18°	—	—	4/20°
Glucinum. See Beryllium.					
Gold —					
chloride, AuCl_3	303.5	—	288°*	dec. 180°	68
Hydrazine, $\text{NH}_2 \cdot \text{NH}_2$	32.05	1.01/15°	1.4	113°	v. soluble
„ hydroxide,					
$\text{N}_2\text{H}_4 \cdot \text{H}_2\text{O}$	50.07	1.030/21°	< -40	119	v. soluble
Hydrobromic acid, HBr	80.93	{ 1.78 A. 2.79 }	-86	-68.7	{ 221/0° 130/100° }
Hydrochloric acid, HCl	36.47	929/0°†	-112.5	-83°-1/755	see p. 124.
Hydrocyanic acid, HCN	27.02	697/18°	-13.8	26.1	∞

* Under chlorine at 1520 mms. † Rupert, 1909. dec. or decomp. = decomposes;
liq. = liquid; r. ht. = red heat; subl. = sublimes; v. = very; ∞ = soluble in all proportions.

INORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.	Solubility in Water.
Hydrofluoric acid, HF . . .	20.01	at./temp. { 988/15° A. 691 }	at./mms. -92°3	at./mms. 19°4	at./temp. 111/35°
Hydriodic acid, HI . . .	127.93	A. 4.38	-51.3	-36°7/752	{ 42,500 V/10° }
Hydrogen —					
peroxide, H ₂ O ₂ . . .	34.02	1.458/0°	-2	80°2/47	v. soluble
selenide, H ₂ Se . . .	81.22	A. 2.805	-64	-42°	331 V/13°
sulphide, H ₂ S . . .	34.08	{ liq. 9 A. 1.178 }	-86	-61.6	{ 305 V/15° see p. 124. }
telluride, H ₂ Te . . .	129.52	A. 4.39	-48	0	soluble
Hydroxylamine, NH ₂ OH . .	33.03	1.227/14°	33°	70°/60	soluble
Iodine —					
trichloride, ICl ₃ . . .	233.3	3.11	101°/16 atm.	dec. 25°	soluble
Iodic acid, HIO ₃ . . .	175.93	4.63/0°	$\frac{1}{2}$ H ₂ O/170°	—	75/16° <i>p.</i>
Iron —					
carbonyl, Fe(CO) ₅ . . .	195.85	{ 1.494/0° A. 6.5 }	-19.7	102°7/764	—
ferrous chloride, FeCl ₂ . .	126.8	2.99/18°	—	volatilizes	50/19°
„ oxide, FeO . . .	71.85	—	—	—	insoluble
„ sulphate, FeSO ₄ ·7H ₂ O . . .	278.03	1.88	64	6 H ₂ O/100°	20.8/10°
„ amm. sulphate, FeSO ₄ (NH ₄) ₂ SO ₄ ·6H ₂ O . . .	392.15	1.81	—	—	{ 18/0° 78/75° }
oxide (magnetic), Fe ₃ O ₄ . .	231.55	5.54	—	—	insoluble
ferric chloride, FeCl ₃ . . .	162.23	{ 2.8/11° A. 11.2/320° }	301	280°-285°	537/100°
„ nitrate, Fe(NO ₃) ₃ ·9H ₂ O .	404.02	1.683/20°	47.2	decomp.	v. soluble
„ oxide, Fe ₂ O ₃ . . .	159.7	5.2-5.3	—	—	insoluble
„ sulphate, Fe ₂ (SO ₄) ₃ (and +9H ₂ O) .	399.91	3.1/18°	—	—	v. slgt. sol.
Lead —					
acetate, Pb(C ₂ H ₃ O ₂) ₂ ·3H ₂ O .	379.2	2.5	3 H ₂ O/75°	280	46/15°
carbonate, PbCO ₃ . . .	267.1	6.4	—	—	decomp.
chloride, PbCl ₂ . . .	277.8	5.8	447°	c. 900	7/0°
iodide, PbI ₂ . . .	460.94	6.12	373	861-954	04/0°
oxide, mon- (litharge), PbO .	223.1	c. 9.3	red heat	—	002/20°
„ red lead, Pb ₃ O ₄ . . .	685.3	9.09/15°	dc. 500°-530°	—	insoluble
„ per- (brown), PbO ₂ . . .	339.1	8.91-9.5	decomp.	—	insoluble
sulphate, PbSO ₄ . . .	303.2	6.23	937°	—	004/18°
Lithium —					
carbonate, Li ₂ CO ₃ . . .	73.88	2.11	618-710	—	see p. 125.
chloride, LiCl . . .	42.40	2.207	491-600	dec. w. ht.	72/0°
nitrate, LiNO ₃ . . .	68.95	2.3-2.4	c. 258	—	35/0°
oxide, Li ₂ O . . .	29.88	2.10/15°	—	—	5/0°
phosphate, Li ₃ PO ₄ ·H ₂ O . .	133.8	2.4/15°	857	—	04
sulphate, Li ₂ SO ₄ . . .	110.0	2.21/15°	818-853	—	26/0°
Magnesium —					
carbonate, MgCO ₃ . . .	84.32	3.0	dec. 350°	—	01
chloride, MgCl ₂ ·6H ₂ O . . .	203.34	1.56/17°	2 H ₂ O/100°	decomp.	54/20°
nitrate, Mg(NO ₃) ₂ ·6H ₂ O . .	256.44	1.46	90°	143	42/18° <i>p.</i>
oxide, MgO . . .	40.32	3.2-3.7	>2000	—	001
phosphate, Mg ₃ (PO ₄) ₂ ·4H ₂ O .	335.2	1.64/15°	—	—	02
sulphate, MgSO ₄ ·7H ₂ O . . .	246.5	1.678/16°	5 H ₂ O/150°	—	27/0°

atm. = atmospheres; dc., dec., or decomp. = decomposes; liq. = liquid; slgt. = slightly; v. = very; w. ht. = white heat.

PHYSICAL CONSTANTS

INORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight ($O = 16$).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.	Solubility in Water.
Manganese—					
carbonate, $MnCO_3$	114.93	at./temp. 3.1–3.7	decomp.	at./mms. —	v. slgt. sol.
chloride, $MnCl_2 \cdot 4H_2O$. .	197.9	1.91	87°6	—	107/10°
nitrate, $Mn(NO_3)_2 \cdot 6H_2O$.	287.05	1.82	87.5	dec. 129°4	54.5/11°p.
oxide, -ous, MnO	70.93	5.1	white heat	—	insoluble
„ -ic, Mn_2O_3	157.86	4.3–4.8	—	—	insoluble
„ tetr-, Mn_3O_4	228.79	4.7–4.9	—	—	insoluble
„ di-, MnO_2	86.93	4.7–5.0	dec. 390	—	insoluble
sulphate,* $MnSO_4 \cdot H_2O$. .	223.06	2.1	18° and 30°†	—	111/54°
Mercury—					
mercurous chloride, $HgCl$.	235.46	{ 6.48 and 7.2 A. 8.21 }	400–500	sublimes	°0002/18°
„ nitrate, $HgNO_3 \cdot 2H_2O$	298.04	4.78	decomp.	—	v. soluble
„ sulphate, Hg_2SO_4	496.07	7.56	melts, dec.	decomp.	°2 cold
mercuric bromide, $HgBr_2$.	359.84	5.7	244	subl. c. 322°	1/9°
„ chloride, $HgCl_2$.	270.92	{ 5.3–5.5 A. 9.8 }	287	303–307	{ 5.4/20°(p) (see p. 125.
„ iodide, red, HgI_2 .	453.84	{ 6.2–6.3 A. 15.6 }	241–257	349	°003/17°
„ „ yellow, HgI_2	453.84	{ 5.9–6.1 A. 15.6 }	241	349	insoluble
„ oxide, HgO	216.0	11.14	dec. r. ht.	—	°005/25°
„ sulphate, $HgSO_4$.	296.07	6.47	dec. r. ht.	—	decomp.
Molybdenum—					
chloride, $MoCl_5$	273.3	A. 9.5/350°	194°	268°	decomp.
oxide, di-, MoO_2	128.0	6.4/10°	—	—	insoluble
„ tri-, MoO_3	144.0	4.4/21°	759	sublimes	°2 cold
Nickel—					
carbonyl, $Ni(CO)_4$	170.7	1.318/17°	–25	43°	insoluble
chloride, $NiCl_2$	129.6	2.56	sublimes	—	35/0°(p)
nitrate, $Ni(NO_3)_2 \cdot 6H_2O$.	290.8	2.06/14°	56°7	136.7	48.5/18°p.
sulphate, $NiSO_4 \cdot 7H_2O$. .	280.86	1.98	98–100	—	31.5/9°
Niobium—					
chloride, penta-, $NbCl_5$. .	270.8	{ 4.4–4.5 A. 9.6/360° }	194	240.5	decomp.
Nitrogen—					
nitric acid, HNO_3	63.02	1.53/15°	–41.3	dec. 86	∞
nitrous oxide, N_2O	44.02	{ 1.226/–89°4 A. 1.614 }	–102	–89°4/74.1	{ 74V/15° (see p. 124.
nitric „ NO	30.01	{ °0013 A. 1.039 }	–167	–153	{ 5.1V/15° (see p. 124.
nitrogen trioxide, N_2O_3 . .	76.02	1.447/–2°	–111	decomp.	soluble
„ peroxide, NO_2 or N_2O_4	46.01	1.49/0° §	–10.1	26°	soluble
„ pentoxide, N_2O_5 . .	108.02	1.64/18°	30	dec. 45–50	soluble
„ oxychloride, $NOCl$.	65.47	1.416/–12°	–60	–5°6/75.1	decomp.
Osmium—					
oxide, tetr-, OsO_4	254.9	A. 8.89	20	100	soluble
Ozone, O_3	48.00	{ °00214 A. 1.659 }	dec. 270°	–119	v. slgt. sol.
Palladium—					
chloride, $PdCl_2 \cdot 2H_2O$. .	213.65	—	dec. r. ht.	—	soluble

* The ordinary salt; also six other hydrates.

† Stable between temps. given.

‡ Also anhy. and $6H_2O$. § Density, p. 26.

dec. or decomp. = decomposes;

r. ht. = red heat; slgt. = slightly; subl. = sublimes; v. = very; ∞ = soluble in all proportions.

INORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 100.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.	Solubility in Water.
Perchloric acid, HClO_4 . . .	100.47	at./temp. 1.76/22°	at./mms. -35	at./mms. 19°/11	at./temp. soluble
Phosphorus —					
bromide, tri-, PBr_3	270.8	{ 2.92/0° A. 9.706 }	-41°5	175	decomp.
chloride, tri-, PCl_3	137.3	{ 1.612/0° A. 4.875 }	-112	76	"
" penta-, PCl_5	208.3	A. 3.6/296°	148	162	"
fluoride, tri-, PF_3	88.04	A. 3.02	-160	-95	"
oxide, tri-, P_2O_5	220.2	1.94/25°	22.5	173	soluble
" tetr-, P_2O_4	126.1	2.54/23°	>100	c. 180	"
" pent-, P_2O_5	142.1	2.39	subl. r. ht.	—	v. soluble
Phosphine, PH_3	34.06	A. 1.185	-133°	-85	slightly sol.
" liquid, P_2H_4	66.11	1.007-1.016	<-10°	57/735	insoluble
Phosphonium chloride, PH_4Cl	70.53	—	26°	sublimes	decomp.
Platinum —					
chloride, tetra-, PtCl_4 . . .	337.0	—	decomp.	—	v. soluble
Potassium —					
bromide, KBr	119.02	2.76/20°	750°	subl. w. ht.	see p. 125.
carbonate, K_2CO_3	138.2	2.29	c. 880	dec. 810°	89/0°
chlorate, KClO_3	122.56	2.34/17°	370	dec. 400°	3/0°
chloride, KCl	74.56	1.99/15°	c. 770	subl. w. ht.	see p. 125.
chromate, bi-, $\text{K}_2\text{Cr}_2\text{O}_7$. . .	294.2	2.69/4°	400	decomp.	5/0°
cyanide, KCN	65.11	1.52/16°	red heat	red heat	122/103°
ferricyanide, $\text{K}_3\text{Fe}(\text{CN})_6$. .	329.21	1.82/17°	decomp.	—	33/4°
ferrocyanide, $\text{K}_4\text{Fe}(\text{CN})_6 \cdot 3\text{H}_2\text{O}$	422.36	1.85/17°	$3\text{H}_2\text{O}/60-80$	—	28/12°
hydroxide, KOH	56.11	2.04	red heat	subl. w. ht.	see p. 125.
iodate, KIO_3	214.02	3.97/18°	560	—	8/20°
iodide, KI	166.1	{ 3.04/24° A. 5.5/1320° }	614-723	—	{ 127/0° see p. 125
nitrate, KNO_3	101.11	2.14°	c. 345	decomp.	see p. 125.
permanganate, KMnO_4 . . .	158.03	2.70/10°	dec. 240°	—	6.4/15
sulphate, K_2SO_4	174.27	2.66/20°	1070	sublimes	9.2/10°
" acid, KHSO_4	136.18	2.24 *; 2.61 †	200	decomp.	36/0°
sulphocyanate, KCNS . . .	97.18	1.91	161	—	217/20°
Radium —					
bromide, RaBr_2	386.24	—	728	—	soluble
Rubidium —					
carbonate, Rb_2CO_3	230.9	—	837	dec. 740°	v. soluble
chloride, RbCl	120.9	2.2	710	—	84/10°
sulphate, Rb_2SO_4	266.97	3.61	—	—	43/10°
Selenium —					
chloride, Se_2Cl_2	229.32	2.91/17°	—	dec. c. 145	decomp.
oxide, SeO_2	111.2	3.95/15°	sub. c. 260	—	v. soluble
Selenious acid, H_2SeO_3 . . .	129.22	3.91/15°7	decomp.	—	"
Selenic acid, H_2SeO_4 . . .	145.22	2.95/15°	58	260	"
Silicon —					
chloride, tetra-, SiCl_4 . . .	170.14	{ 1.520 A. 5.94 }	-89	57.5	decomp.
fluoride, SiF_4	104.3	A. 3.57	-102	-107	"

* Monoclinic.

† Rhombic.

amorph. = amorphous; cryst. = crystalline; dec. or decomp. = decomposes; r. ht. = red heat; sub. or subl. = sublimes; v. = very; w. ht. = white heat.

For general heading, see p. 109.

* Practically same for ordinary table salt as for pure salt (Härker).
 anhy. = anhydrous ; dec. or decomp. = decomposes ; hydr. = hydrated ; r. ht. = red heat ;
 w. ht. = white heat ; ∞ = soluble in all proportions.

INORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.	Solubility in Water.
Thallium —		at./temp.	at./mms.	at./mms.	at./temp.
carbonate, Tl_2CO_3	468.0	7.1	272°	decomp.	4/15°
chloride, tri-, TlCl_3	310.38	—	25	"	v. soluble
oxide (thallous), Tl_2O	424.0	—	300	—	v. soluble
sulphate, Tl_2SO_4	504.07	6.77	632	decomp.	4.7/15°
Thorium —					
nitrate, $\text{Th}(\text{NO}_3)_4 \cdot 12\text{H}_2\text{O}$.	696.2	—	—	—	v. soluble
oxide, ThO_2	264.0	9.87/15°	infusible	—	insoluble
Tin —					
chloride (stannous), SnCl_2 . .	189.92	—	249°	620°	270/15°
" (stannic), SnCl_4	260.84	{ 2.27/20° } A. 9.2	— 33	114.1	soluble
oxide (stannous), SnO	135.0	6.3	dec. r. ht.	—	insoluble
" (stannic), SnO_2	151.0	6.6-6.9	1130	—	"
Titanium —					
chloride, tetra-, TiCl_4	189.94	{ 1.76/0° } A. 6.836	— 25	136	decomp.
oxide, di-, TiO_2	80.1	3.7-4.2	c. 1500	—	insoluble
Tungsten —					
chloride, hexa-, WCl_6	396.76	A. 13.3/350°	275	347	"
oxide, tri-, WO_3	232.0	7.2	red heat	—	"
Uranium					
oxide, di-, UO_2	270.5	10.9	oxidises	—	"
" (green), U_3O_8	843.5	7.3	decomp.	—	—
" (yellow), UO_3	286.5	5.1	decomp.	—	—
" (black), U_3O_8	557.0	8.4-9.2	—	—	—
Uranyl chloride, UO_2Cl_2 . . .	341.42	—	fusible	decomp.	320/18°
" nitrate, $\text{UO}_2(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$	502.62	2.81	59°5	118°	200
Vanadium —					
chloride, tetra-, VCl_4	192.9	{ 1.86 } A. 6.69	— 18	154	soluble
oxide, pent-, V_2O_5	182.1	3.5/20°	658	—	0.8/20°
Zinc —					
carbonate, ZnCO_3	125.37	4.4	dec. 300°	—	0.001/15°
chloride, ZnCl_2	136.29	2.91/25°	262°?	730	330/10°
sulphate, $\text{ZnSO}_4 \cdot 7\text{H}_2\text{O}$. . .	287.55	{ 1.96 } 3.4 anhy.	6H ₂ O/100°	{ 7H ₂ O at } { red heat. }	{ 42/0° } 80.8/100°
sulphide, ZnS	97.44	4.0	1050°	—	insoluble
Zirconium —					
oxide, ZrO_2	122.6	5.1-5.7	infusible	—	"

anhy. = anhydrous; dec. or decomp. = decomposes; r. ht. = red heat; v. = very.

FREEZING MIXTURES

Parts by weight.	Temp.	Parts by weight.	Temp.
1 of NH_4NO_3 , 1 of water . . .	— 15° C.	2 of snow or crushed ice, 1 of NaCl	— 18°
8 of Na_2SO_4 , 5 of water . . .	— 17	3 of snow, 4 of cryst. CaCl_2 .	— 48

PHYSICAL CONSTANTS

ORGANIC COMPOUNDS

Formula (Molecular) Weight, Density, Melting and Boiling Points.

For general heading, see p. 109.

Substance and Formula.	Formula weight (0 = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.
Acetaldehyde, $\text{CH}_3 \cdot \text{CHO}$	44.03	at./temp. 788/16° C.	at./mms. -120°	at./mms. 20° 8
Acetic acid, $\text{CH}_3 \cdot \text{COOH}$	60.03	1.05/20°	16.7	118.5, Y.
Aceto-acetic ether, $\text{CH}_3\text{CO} \cdot \text{CH}_2\text{CO}_2$ C_2H_5	130.1	1.028/20°	< -80	181
Acetone, CH_3COCH_3	58.05	797/15°	-95	56.5
Acetylene, C_2H_2	26.02	{46/-7°} A. .91	-81.5/89.5*	-85
Acrylic acid, $\text{CH}_2 : \text{CHCO}_2\text{H}$	72.03	1.062/16°	10	140
Alizarine, $\text{C}_6\text{H}_4(\text{CO})_2\text{C}_6\text{H}_2(\text{OH})_2$	240.1	—	290	430
Allyl alcohol, $\text{CH}_2 : \text{CH} \cdot \text{CH}_2\text{OH}$	58.05	.85/15°	liquid	96.7
" chloride, $\text{CH}_2 : \text{CHCH}_2\text{Cl}$	76.46	.937/19°	liquid	46
" thiocyanate, $\text{CH}_2 : \text{CHCH}_2\text{CNS}$	99.08	1.017/10°	liquid	151
Amyl acetate, $\text{C}_5\text{H}_{11} \cdot \text{CH}_3\text{CO}_2$	130.1	.879/20°	liquid	148
" alcohol (n.), $\text{CH}_3(\text{CH}_2)_4\text{CH}_2\text{OH}$	88.10	.812/20°	liquid	137
" " (act.), $\text{CH}_3\text{C}_2\text{H}_5\text{CHCH}_2$ - OH	88.10	.825/0°	liquid	129
" " (sec.), $\text{C}_3\text{H}_7\text{CH}(\text{OH})\text{CH}_3$	88.10	.825/0°	liquid	118.5/753
" " (tert.), $(\text{CH}_3)_2\text{C}(\text{OH})$ - C_2H_5	88.10	.814/15°	-12°	102.5
Aniline, $\text{C}_6\text{H}_5 \cdot \text{NH}_2$	93.07	1.023/15°	-8	183.9
Anisol, $\text{C}_6\text{H}_5\text{OCH}_3$	108.1	.99/25°	-37.8	155
Anthracene, $\text{C}_6\text{H}_4 : \text{C}_6\text{H}_2\text{C}_6\text{H}_4$	178.1	1.15	216	351
Antimony trimethyl, $\text{Sb}(\text{CH}_3)_3$	165.3	1.52/15°	liquid	86
Asparagine(l.), $\text{C}_4\text{H}_7\text{NH}_2\text{CO}_2\text{H} \cdot \text{CONH}_2$	132.1	1.55/4°	decomp.	decomp.
Benzaldehyde, $\text{C}_6\text{H}_5\text{CHO}$	106.1	1.05/15°	-13° 5	179.5
Benzene, C_6H_6	78.05	.879/20°	5.4	80.2, Y.
Benzoic acid, $\text{C}_6\text{H}_5 \cdot \text{COOH}$	122.0	1.20/21°	121.4	249.2
Benzophenone, $(\text{C}_6\text{H}_5)_2\text{CO}$	182.1	1.098/50°	48	306
Benzoyl chloride, $\text{C}_6\text{H}_5\text{COCl}$	140.5	1.212/20°	-1	198/749
Benzyl alcohol, $\text{C}_6\text{H}_5\text{CH}_2\text{OH}$	108.1	1.043/20°	liquid	206.5
Beryllium ethyl, $\text{Be}(\text{C}_2\text{H}_5)_2$	67.18	—	—	187
Bismuth triethyl, $\text{Bi}(\text{C}_2\text{H}_5)_3$	295.1	2.3/18°	—	107
Borneol (i.), $\text{C}_{10}\text{H}_{17}\text{OH}$	154.1	1.01	210	sublimes
Bromo benzene, $\text{C}_6\text{H}_5\text{Br}$	157.0	1.49/20°	-31.1	156, Y.
Butyl alcohol (n.), $\text{CH}_3(\text{CH}_2)_3\text{CH}_2 \cdot \text{OH}$	74.08	.81/20°	liquid	117.5
" (sec.), $\text{CH}_3\text{CHOH} \cdot \text{C}_3\text{H}_7$	74.08	.819/22°	—	99.8
" carbinol (tert.), $(\text{CH}_3)_3\text{C} \cdot \text{CH}_2\text{OH}$	88.10	.812/20°	52	113
" chloride, $\text{CH}_3(\text{CH}_2)_3\text{Cl}$	92.53	.887/20°	liquid	78
" ether, $(\text{C}_4\text{H}_9)_2\text{O}$	130.1	.77/20°	—	141
Butyric acid (n.), $\text{CH}_3(\text{CH}_2)_3\text{COOH}$	88.06	.96/19°	-8	162.3
" " (iso), $(\text{CH}_3)_2\text{CHCOOH}$	88.06	.950/20°	-79	155
Calcodylic acid, $(\text{CH}_3)_2\text{AsO} \cdot \text{OH}$	138.0	—	200	—
Caffeine, $\text{C}_8\text{H}_{10}\text{N}_4\text{O}_2 \cdot \text{H}_2\text{O}$	212.3	1.23/19°	234	sublimes
Camphor, $\text{C}_{10}\text{H}_{16}\text{O}$	152.1	.992/10°	176.4	205.3
Camphoric acid (d.), $\text{C}_8\text{H}_{14}(\text{COOH})_2$	200.1	1.19	178	decomp.
Caproic acid, $\text{CH}_3(\text{CH}_2)_4\text{COOH}$	116.1	.929/20°	8	205
Carbolic acid. See Phenol.				
Carbon bisulphide, CS_2	76.14	1.292/0°	-110	46.2
" oxysulphide, COS	60.07	2.104	—	gas
" tetrachloride, CCl_4	153.8	1.582/21°	-30	76.7, Y.

* Mackintosh, 1907; decomp. = decomposes; l., = levo-rotatory (see p. 78). Y., Young, *Journal de Phys.*, Jan., 1909.

ORGANIC COMPOUNDS (contd.)

For general heading, see p. 109.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, ° C.	Boiling Point, ° C.
		at./temp.	at./mms.	at./mms.
Cellulose, (C ₆ H ₁₀ O ₅) _x	162.1	1.525	—	—
Chlor acetic acid, CClH ₂ :COOH . .	94.48	1.39/75°	63°	186°
„ benzene, C ₆ H ₅ Cl	112.5	1.118/10°	-40	132, Y.
Chloral hydrate, CCl ₃ .CH(OH) ₂ . .	165.4	1.9	-57	97.5
Chloroform, CHCl ₃	119.4	1.526/0°	-70	61.2
Chrysene, C ₁₈ H ₁₂	228.1	—	250	sublimes
Cineol, C ₁₀ H ₁₈ O	154.2	.92	-1	176
Cinnamic acid, C ₆ H ₅ CH:CHCOOH	148.1	1.247	133	300
„ aldehyde, C ₆ H ₅ CH:CH-CHO	132.1	1.05/24°	-7.5	—
Citric acid, (CO ₂ HCH ₂) ₂ C(OH)CO ₂ H + H ₂ O	192.1	1.54	153	decomp.
Collidine, α CH ₃ .C ₆ H ₃ N.C ₂ H ₅ . .	121.1	.953/22°	—	180
Conine (d.), 1:2, C ₈ H ₁₀ N.C ₃ H ₇ . .	127.2	.849/25°	-2.5	170
Cresol (o.), CH ₃ C ₆ H ₄ OH	108.1	1.005	30	191
Cyanic acid, HCNO	43.02	1.14/0°	liquid	dec. o
Cyanogen, C ₂ N ₂	52.02	{liq. .866/17° A. 1.806}	-35	-20.7
Cymene (p.), CH ₃ .C ₆ H ₄ .C ₃ H ₇ . .	134.12	.852/25°	liquid	175
Dextrin, C ₁₂ H ₂₀ O ₁₀	324.2	1.04	—	—
Diacetyl, CH ₃ CO.COCH ₃	86.05	.973	—	87.7
Dichlor acetic acid, CHCl ₂ .COOH .	128.9	1.522/15°	-4	190
Diethyl amine, (C ₂ H ₅) ₂ NH	73.13	.706/20°	-40	55.5
„ aniline, (C ₂ H ₅)NC ₆ H ₅	149.2	.94/18°	liquid	213.5
„ ketone, C ₂ H ₅ COC ₂ H ₅	86.08	.83/0°	—	103
Dimethyl amine, (CH ₃) ₂ NH	45.07	.686/-6°	liquid	8 to 9
„ tartrate, (CH ₃) ₂ C ₄ H ₄ O ₆	178.1	1.341/15°	48	280
Dinitrobenzene (m.), C ₆ H ₄ (NO ₂) ₂ .	168.1	1.37	91	297
Diphenyl, C ₆ H ₅ .C ₆ H ₅	154.1	1.16	70.5	235
Diphenylamine, (C ₆ H ₅) ₂ NH	169.1	1.159	54	310
Epichlorhydrine, C ₃ H ₅ ClO	92.49	1.203/0°	—	116
Erythrite, (CH ₂ OH.CHOH) ₂	122.1	1.45/17°	112	330
Ethane, CH ₃ .CH ₃	30.05	{liq. .446/0° A. 1.036}	-171.4	-85.4/749
Ether, C ₂ H ₅ OC ₂ H ₅	74.08	.718/17°	-117	34.6, Y.
Ethyl acetate, CH ₃ CO ₂ .C ₂ H ₅ . . .	88.06	.903/18°.5	-83.8	77.1
„ aceto-acetate, CH ₃ COCH ₂ CO ₂ .C ₂ H ₅	130.1	1.028/20°	<-80	181
„ alcohol, C ₂ H ₅ OH	46.05	.7937/15°	-112.3	78.3, Y.
„ amine, C ₂ H ₅ H ₂ N	45.07	.699/8°	-85	187
„ benzoate, C ₆ H ₅ CO ₂ .C ₂ H ₅	150.1	1.05/16°	111-116	211.2
„ bromide, C ₂ H ₅ .Br	108.96	1.45/15°	-116	38.4
„ butyrate, C ₃ H ₇ .COOC ₂ H ₅	116.1	.898/18°	—	120
„ chloride, C ₂ H ₅ Cl	64.50	{.921/0° A. 2.219}	liquid	12.5
„ cyanide, C ₂ H ₅ .CN	55.05	.794/7°	-103	97
„ formate, HCOOC ₂ H ₅	74.05	.938/0°	—	54.3, Y.
„ iodide, C ₂ H ₅ I	156.0	1.944/14°	liquid	72.3
„ isobutyrate, (CH ₃) ₂ CHCOOC ₂ H ₅	116.1	.890/0°	—	110.1
„ mercaptan, C ₂ H ₅ SH	62.11	.839/20°	-22	36.2
„ nitrate, C ₂ H ₅ NO ₃	91.08	1.116/15°	-112	87

dec. or decomp. = decomposes.

Y., Young, *Journ. de Phys.*, Jan., 1909.

PHYSICAL CONSTANTS

ORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight (0 = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.
		at./temp.	at./mms.	at./mms.
Ethyl propionate, $C_2H_5CO_2C_2H_5$. . .	102.1	896/16°	—	99°0
„ salicylate, $C_6H_4(HO)CO_2C_2H_5$. . .	166.1	1.184/20°	—	231.5
„ sulphide, $(C_2H_5)_2S$. . .	90.15	837/20°	liquid	92.6
„ tartrate (d.), $C_4H_4O_6(C_2H_5)_2$. . .	206.1	1.206/20°	—	280
„ valerate, $C_4H_9CO_2C_2H_5$. . .	130.1	876/20°	—	144.5
Ethylene, $CH_2 : CH_2$	28.03	{liq. 61 } {A. 9784 }	— 169	— 102.7
„ bromide, di-, $CH_2Br \cdot CH_2Br$. . .	187.9	2.19/11°	9.5	131.6
„ chloride, di-, $CH_2Cl \cdot CH_2Cl$. . .	98.93	1.28/0°	— 40	83.7
„ oxide, $<(CH_2)_2O$	44.03	897/0°	liquid	135.746
Ethylidene chloride, $CH_3 \cdot CHCl_2$. . .	98.93	1.186/12°	liquid	59.9
Eucalyptol, $C_{10}H_{18}O$	154.1	927/20°	— 1	176
Eugenol, $C_6H_3 \cdot (OH) \cdot OCH_3 \cdot C_2H_5$. . .	164.1	1.0779/0°	liquid	247.5
Fluor benzene, C_6H_5F	96.04	1.024/20°	40°	85.2, Y.
Formic acid, $H \cdot COOH$	46.02	1.22/20°	8.6	100.8
Formaldehyde, $H \cdot COH$	30.02	{815/— 20° } {A. 1.6 }	—	— 21
Fructose (d.), $CH_2OH[CHOH]_2CO \cdot CH_2OH$	180.1	1.55/0°	95	—
Fumaric acid, $(COOH \cdot CH)_2$	116.0	1.625	286	—
Furfural, $C_4H_3O \cdot COH$	96.03	1.159/20°	liquid	161
Galactose (d.), $CHO[CHOH]_4CH_2OH$. . .	180.1	—	163	—
Glucose (d.), $CHO(HCOH)_4CH_2OH$. . .	198.1	1.54—1.57	146	—
Glutaric acid, $COOH(CH_2)_3COOH$. . .	132.1	—	91	299
Glycerine, $OHCH_2 \cdot CHOH \cdot CH_2OH$. . .	92.06	1.26/20°	17	290
Glycocoll, CH_2NH_2COOH	75.08	1.161	G. 234	—
Glycol, $CH_2OH \cdot CH_2OH$	62.05	1.125/25°	— 17.4	197.4
Glycollic acid, $CH_2OH \cdot COOH$	76.03	—	78	decomp.
Glyoxal, $CHO \cdot CHO$	58.02	—	—	dec. 160
Glyoxalic acid, $CHO \cdot COOH + H_2O$. . .	92.03	syrup	—	with steam
Grape sugar. <i>See</i> Glucose.				
Heptane (n.), $CH_3(CH_2)_5CH_3$	100.1	688/15°	—	98.4, Y.
Hexane (n.), $CH_3(CH_2)_4CH_3$	86.12	658/21°	liquid	69, Y.
„ di-isopropyl, $[(CH_3)_2CH]_2$	86.12	668/17°	liquid	58.1, Y.
Hydrocyanic acid, HCN	27.05	697/18°	— 14	26.1
Indigo, $C_6H_4 \begin{smallmatrix} CO \\ \diagup \\ NH \end{smallmatrix} > C : C < \begin{smallmatrix} CO \\ \diagdown \\ NH \end{smallmatrix} > C_6H_4$	262.2	1.35	—	subl. 156°
Indol, $C_8H_7NHCH : CH$	117.1	—	52	245
Iodoform, CHI_3	393.8	2.25/25°	119	subl. & dec.
Isatine, $C_8H_4 \begin{smallmatrix} CO \\ \diagup \\ N \end{smallmatrix} > COH$	147.1	—	201	sublimes
Isoamyl acetate, $CH_3 \cdot COOC_5H_{11}$. . .	130.1	876/15°	—	140
„ alcohol, $(CH_3)_2CH(CH_2)_2OH$. . .	88.10	81/20°	— 134	129.7
Isobutane, $(CH_3)_2CHCH_3$	58.08	—	—	116.3
Isobutyl alcohol, $(CH_3)_2CH \cdot CH_2OH$. . .	74.08	800/18°	liquid	108.4
„ amine, $(CH_3)_2CHCH_2NH_2$	73.13	736/15°	—	68
Isobutyric acid, $(CH_3)_2CH \cdot COOH$. . .	88.06	949/20°	— 79	155.5
Isopentane, $(CH_3)_2CHCH_2CH_3$	72.10	628/14°	—	27.9
Isopropyl acetate, $CH_3COOCH(CH_3)_2$. . .	102.1	917	—	90—93
„ alcohol, $(CH_3)_2CH(OH)$	60.06	789/20°	liquid	82.8

d., dextro-rotatory (see p. 78); dec. or decomp. = decomposes; subl. = sublimes; Y., Young,
Journ. de Phys., Jan., 1909.

ORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.
Isopropyl amine, $(\text{CH}_3)_2\text{CHNH}_2$. . .	59.11	at./temp. .690/18°	at./mms, liquid	at./mms, 31°5/743
„ cyanide, $(\text{CH}_3)_2\text{CHCN}$. . .	69.07	—	liquid	107-108
Isoquinoline, $\text{C}_8\text{H}_4\text{C}_3\text{H}_3\text{N}$. . .	129.1	1.098/20°	24.6	240
Isovaleric acid, $(\text{CH}_3)_2\text{CHCH}_2\text{COOH}$. . .	102.1	.931/20°	-51	176.3
Lactic acid (i.), $\text{CH}_3\text{CHOH}\cdot\text{COOH}$. . .	90.05	1.248/15°	—	83/1 mm.
Lactose. <i>See</i> Milk sugar.				
Maleic acid, $(\text{COOH}\cdot\text{CH})_2$. . .	116.0	1.59	100	decomp.
Malic acid (i.), $\text{COOH}\cdot\text{CHOH}\cdot\text{CH}_2\cdot\text{COOH}$. . .	134.0	1.60/20°	130-1	—
Malonic acid, $\text{COOH}\cdot\text{CH}_2\cdot\text{COOH}$. . .	104.0	—	132	decomp.
Maltose, $\text{C}_{12}\text{H}_{22}\text{O}_{11} + \text{H}_2\text{O}$. . .	360.2	1.54/17°	—	—
Mercury methyl, $(\text{CH}_3)_2\text{Hg}$. . .	230.0	3.07	liquid	96
Mesitylene, 1 : 3 : 5, $\text{C}_6\text{H}_3(\text{CH}_3)_3$. . .	120.1	.869/10°	—	164.5
Methane, CH_4 . . .	16.03	liq. .416/-164°	-184	-164
Methyl alcohol, CH_3OH . . .	32.03	.796/15°	-94.9	64.7, Y.
„ acetate, $\text{CH}_3\text{COO}\cdot\text{CH}_3$. . .	74.05	.941/14°	-101.2	57.1
„ amine, $\text{CH}_3\text{H}_2\text{N}$. . .	31.08	{.699/-11° A 1.08}	gas	-6.7/756
„ borate, $(\text{CH}_3)_3\text{BO}_3$. . .	104.1	.94/0°	—	65
„ chloride, CH_3Cl . . .	50.48	{.920/18° A 1.73}	—	-24.1
„ ether, $(\text{CH}_3)_2\text{O}$. . .	46.05	A 1.62	gas	-23.6
„ ethyl ether, $\text{CH}_3\cdot\text{O}\cdot\text{C}_2\text{H}_5$. . .	60.06	.725/0°	—	10.8
„ formate, $\text{HCOO}\cdot\text{CH}_3$. . .	60.03	.986/11°	—	31.9, Y.
„ iodide, CH_3I . . .	142.0	2.285/15°	liquid	42.3
„ isobutyrate, $(\text{CH}_3)_2\text{CHCOOCH}_3$. . .	102.1	.912/0°	—	92.3
„ mercaptan, $\text{CH}_3\cdot\text{SH}$. . .	48.09	—	—	5.8/752
„ nitrate, $\text{CH}_3\cdot\text{NO}_3$. . .	77.03	1.217/15°	liquid	65 explodes
„ nitrite, $\text{CH}_3\cdot\text{NO}_2$. . .	61.03	.991/15°	—	-12
„ phosphine, $\text{CH}_3\text{H}_2\text{P}$. . .	48.04	—	gas	-14
„ propionate, $\text{C}_2\text{H}_5\text{COO}\cdot\text{CH}_3$. . .	88.06	.937/0°	—	79.7
„ salicylate, $\text{C}_6\text{H}_4(\text{OH})\text{COOCH}_3$. . .	152.1	1.182/15°	-30	224
„ sulphide, $(\text{CH}_3)_2\text{S}$. . .	62.12	.845/21°	liquid	c. 38
Methylene bromide, CH_2Br_2 . . .	173.9	2.493	—	98.5
Milk sugar, $\text{C}_{12}\text{H}_{22}\text{O}_{11} + \text{H}_2\text{O}$. . .	360.2	1.525/20°	203 dec.	decomp.
Morphine, $\text{C}_{17}\text{H}_{19}\text{NO}_3 + \text{H}_2\text{O}$. . .	303.2	1.32	—	decomp.
Naphthalene, C_{10}H_8 . . .	128.1	1.152/15°	80	218.1
Naphthol (α), $\text{C}_{10}\text{H}_7\text{OH}$. . .	144.1	1.224/4°	95	c. 279
Naphthyl amine (α), $\text{C}_{10}\text{H}_7\text{H}_2\text{N}$. . .	143.1	—	50	300
Nicotine (l.), $\text{C}_{10}\text{H}_{14}\text{N}_2$. . .	162.2	1.01/20°	dec. 250°	246.7/745
Nitro benzene, $\text{C}_6\text{H}_5\text{NO}_2$. . .	123.1	1.187/14°	3.6	209.4/745
„ ethane, $\text{C}_2\text{H}_5\text{NO}_2$. . .	75.08	1.056	194-196	114.4
„ methane, CH_3NO_2 . . .	61.07	1.144/15°	liquid	101.7
Octane (n.), $\text{CH}_3(\text{CH}_2)_6\text{CH}_3$. . .	114.1	.719/0°	liquid	125.8, Y.
Oleic acid, $\text{CH}_3(\text{CH}_2)_7\text{CH}:\text{CH}(\text{CH}_2)_7\cdot\text{COOH}$. . .	282.3	.891/12°	14	286/100
Palmitic acid, $\text{CH}_3(\text{CH}_2)_{14}\text{COOH}$. . .	256.3	.846/7.6°	62.6	278/100
Paraldehyde, $(\text{CH}_3\cdot\text{HCO})_3$. . .	132.1	.994/20°	10.5	124
Penta methylene, $(\text{CH}_2)_5$. . .	70.08	.751/20°	—	50.6
„ diamine (cadaverine), „ $\text{NH}_2(\text{CH}_2)_5\text{NH}_2$. . .	102.2	.917/0°	—	178

dec. or decomp. = decomposes; l., laevo-rotatory (see p. 78); Y., Young, *Journ. de Phys.*, Jan., 1909.

PHYSICAL CONSTANTS

ORGANIC COMPOUNDS (contd.)

For general heading, see p. 109.

Substance and Formula.	Formula weight (0 = 16).	Density, gms./c.c.	Melting Point, °C.	Boiling Point, °C.
Pentane (n.), $\text{CH}_3(\text{CH}_2)_3\text{CH}_3$	72.10	at./temp. .634/15°	at./mms. -200	at./mms. 36°2, Y.
Phenetol, $\text{C}_6\text{H}_5\text{OC}_2\text{H}_5$	122.08	.963/25°	-34	171
Phenol, $\text{C}_6\text{H}_5\cdot\text{OH}$	94.05	1.06/33°	42.7	181.5
Phenyl acetic acid, $\text{C}_6\text{H}_5\text{CH}_2\text{COOH}$	136.1	1.23	76.5	265
„ cyanide, $\text{C}_6\text{H}_5\text{CN}$	103.1	1.008/17°	-17	190
„ hydrazine, $\text{C}_6\text{H}_5\text{HN}\cdot\text{NH}_2$	108.1	1.1/23°	23	233
Phloroglucin, 1:3:5, $\text{C}_6\text{H}_3(\text{OH})_3\cdot 2\text{H}_2\text{O}$	162.1	—	218 anhy.	sublimes
Phthalic acid, o. $\text{C}_6\text{H}_4(\text{COOH})_2$	166.1	1.59	180-200	—
„ anhydride, $\text{C}_6\text{H}_4<(\text{CO})_2>\text{O}$	148.0	1.53/4°	128	284
Picoline (a), $\text{CH}_3\cdot\text{C}_6\text{H}_4\text{N}$	93.07	.933/22°	liquid	129
Picric acid, 1:2:4:6, $\text{C}_6\text{H}_2\text{OH}(\text{NO}_2)_3$	229.1	1.813	122.5	explodes
Propane, $\text{CH}_3\cdot\text{CH}_2\cdot\text{CH}_3$	44.07	.535	-195	-(38-39)
Propionic acid, $\text{CH}_3\cdot\text{CH}_2\cdot\text{COOH}$	74.05	.995/20°	-22	140
Propyl acetate (n.), $\text{CH}_3\text{COO}\cdot\text{C}_3\text{H}_7$	102.0	.891/18°	liquid	101.6
„ alcohol (n.), $\text{CH}_3\text{CH}_2\text{CH}_2\cdot\text{OH}$	60.06	.804/20°	—	97.2
„ chloride (n.), $\text{CH}_3\text{CH}_2\text{CH}_2\text{Cl}$	78.51	.891/18°	—	46.5
„ formate, $\text{H}\cdot\text{COO}\cdot\text{C}_3\text{H}_7$	88.06	.909/17°	—	80.9, Y.
„ iodide, $\text{CH}_3\cdot\text{CH}_2\cdot\text{CH}_2\text{I}$	170.0	1.745/20°	—	102
Propylene, $\text{CH}_3\cdot\text{CH}:\text{CH}_2$	42.05	A. 1.498	gas	-50.2
Pseudo-cumene, 1:2:4, $\text{C}_6\text{H}_3(\text{CH}_3)_3$	120.1	.879/20°	—	169.8
Pyridine, $\text{C}_5\text{H}_5\text{N}$	79.08	.985/15°	liquid	117
Pyrogallol (—ic acid, or “pyro”), 1:2:3, $\text{C}_6\text{H}_3(\text{OH})_3$	126.1	1.46/40°	133	293
Pyrrol, $(\text{CH}_2)_4>\text{NH}$	67.08	.967/21°	liquid	131
Quinoline, $\text{C}_6\text{H}_4<\begin{smallmatrix} \text{CH}\cdot\text{CH} \\ \text{N}\cdot\text{CH} \end{smallmatrix}>$	129.1	1.094/20°	19.5	241
Quinine, $\text{C}_{20}\text{H}_{24}\text{N}_2\text{O}_2$	324.3	—	174.9	—
„ sulphate, $(\text{C}_{20}\text{H}_{24}\text{N}_2\text{O}_2)_2\cdot$ $\text{H}_2\text{SO}_4 + 7\text{H}_2\text{O}$	872.7	—	205, dry	—
Racemic acid, $(\text{COOH}\cdot\text{CH}(\text{OH}))_2\cdot$ H_2O	168.1	1.69/7°	205	—
Rochelle salt (d.), $\text{KNaC}_4\text{H}_4\text{O}_6$	—	—	—	—
Rosaniline (p.), $(\text{C}_6\text{H}_4\text{NH}_2)_3\text{COH}$	305.2	—	—	—
Saccharin, $\text{C}_6\text{H}_4<\text{COSO}_2>\text{NH}$	183.1	—	220 dec.	—
Salicylic acid, $\text{OH}\cdot\text{C}_6\text{H}_4\cdot\text{COOH}$	138.0	1.48/4°	158	sublimes
Sodium ethyl, NaC_2H_5	52.04	—	—	—
Stearic acid, $\text{CH}_3(\text{CH}_2)_{16}\text{COOH}$	284.3	.843/80°	69.3	291/100
Stearine, $(\text{C}_{18}\text{H}_{36}\text{O}_2)_3\cdot\text{C}_3\text{H}_5$	890.9	.924/65°	—	—
Succinic acid, $\text{COOH}(\text{CH}_2)_2\text{COOH}$	118.0	1.55	185	235
Sugar, cane-, $\text{C}_{12}\text{H}_{22}\text{O}_{11}$	342.2	1.588/20°	—	185
Sulphanilic acid (p.), $\text{NH}_2\cdot\text{C}_6\text{H}_4\cdot\text{SO}_3\text{H}$ $\cdot 2\text{H}_2\text{O}$	209.2	—	chars	—
Sulphonal, $(\text{CH}_3)_2\text{C}(\text{SO}_2\text{C}_2\text{H}_5)_2$	228.2	—	125	300 dec.
Tartaric acid (l. or meso), $\text{COOH}\cdot$ $[\text{CHOH}]_2\text{COOH}\cdot\text{H}_2\text{O}$	168.1	1.67	142 anhy.	—
„ „ (d.), $\text{COOH}(\text{CHOH})_2\cdot$ COOH	150.0	1.76/7° P.	170	—
„ „ (l.), $\text{COOH}(\text{CHOH})_2\cdot$ COOH	150.0	1.76	170	—
Terephthalic acid (p.), $\text{C}_6\text{H}_4(\text{COOH})_2$	166.0	—	—	sublimes
Terpenol, $\text{C}_{10}\text{H}_{18}\text{O}$	154.1	—	70	—

anhy. = anhydrous; d. = dextro-rotatory (see p. 78); P., Perkin; dec. = decomposes;
l., lævo-rotatory (see p. 78); Y., Young.

ORGANIC COMPOUNDS (*contd.*)

For general heading, see p. 109.

Substance and Formula.	Formula weight (O = 16).	Density, gms./c.c.	Melting Point, ° C.	Boiling Point, ° C.
Terpineol, $C_{10}H_{17}HO$	154.1	at./temp. .936/20°	at./mms. 35°	at./mms. 218°
Tetrabromethylene, $CBr_2 \cdot CBr_2$	343.8	—	53	—
Theobromine, $C_7H_8N_4O_2$	180.2	—	330	decomp.
Thiocyanic acid, $HCNS$	59.09	—	-12.5	200 dec.
Thiourea, $NH_2 \cdot CS \cdot NH_2$	76.12	1.42	180	—
Thymol, 3 : 2 : 1, $(CH_3)_2 : \dot{C}H \cdot C_6H_3 -$ $(CH_3)OH$	150.1	.994/0°	50	232
Tin tetramethyl, $Sn(CH_3)_4$	179.1	1.314	—	78
Toluene, $C_6H_5 \cdot CH_3$	92.06	.866/20°	-97	111
Toluidine (o.), $CH_3C_6H_4 \cdot NH_2$	107.1	.999/20°	liquid	197
" (p.), $CH_3C_6H_4 \cdot NH_2$	107.1	1.046/—	45	108
Trichloroacetic acid, $CCl_3 \cdot COOH$	163.4	1.63/61°	52.3	195
Triethyl amine, $(C_2H_5)_3N$	101.2	.735/15°	liquid	89
" arsine, $(C_2H_5)_3As$	162.1	1.15/17°	liquid	{ 140/736 dec.
" phosphine, $(C_2H_5)_3P$	118.1	.812/15°	liquid	127/744
Trimethyl amine, $(CH_3)_3N$	59.08	.673/0°	—	3.5
" arsine, $(CH_3)_3As$	120.0	—	—	<100
" bismuth, $(CH_3)_3Bi$	253.1	2.30/18°	—	110
" carbinol, $(CH_3)_3C \cdot OH$	74.08	.786/20°	25	82.9
" phosphine, $(CH_3)_3P$	76.07	>1	liquid	41
Trinitro benzene (s.), 1 : 3 : 5, $C_6H_3 -$ $(NO_2)_3$	213.1	—	121.2	decomp.
Turpentine (pinene), $C_{10}H_{16}$	136.1	.865/15°	—	159
Urea, $NH_2 \cdot CO \cdot NH_2$	60.11	1.32	132	decomp.
Valeric acid (n.), $CH_3(CH_2)_3 \cdot COOH$	102.1	.943/20°	-58.5	186.4
Xylene (o.), $C_6H_4(CH_3)_2$	106.1	.756/14°	-28	142
" (m), "	106.1	.878/0°	-54	139.8
" (p), "	106.1	.862/20°	15	138
Zinc ethyl, $Zn(C_2H_5)_2$	123.5	1.182/18°	-28	118
" methyl, $Zn(CH_3)_2$	95.42	1.386/10°	-40	46

dec. or decomp. = decomposes.

ELECTROCHEMICAL EQUIVALENTS

Faraday's laws of electrolysis are expressed by $m = izt$, where m is the mass in grammes of an ion liberated in t secs. by a current of i amperes; z is the electrochemical equivalent of the ion, *i.e.* the mass liberated by 1 ampere in 1 second.

The exactness of Faraday's laws is obscured in many cases by secondary chemical reactions, and the values of the different electrochemical equivalents are practically always derived by calculation from that of silver, which has been accurately determined (see p. 8). Electrochemical equivalents are proportional to chemical equivalents.

atomic weight of element
Chemical equivalent = —————
valency of element for electrolyte used

Element.	Chemical equivalent.	z.
Silver	107.88/1	0.0011183 gm. sec. ⁻¹ amp. ⁻¹
Copper	63.57/2	0.0003295 " " "
Hydrogen	1.008/1	0.00001045 " " (see p. 106)

SOLUBILITIES

SOLUBILITIES OF GASES IN WATER

AIR IN WATER

1000 c.c.s. of water saturated with air at a pressure of 760 mms. contain the following volumes of dissolved oxygen, etc., in c.c.s. at 0° and 760 mms.

	Temperature of Water.						
	0° C.	5°	10°	15°	20°	25°	30°
	c.c.s.						
Oxygen	10.19	8.9	7.9	7.0	6.4	5.8	5.3
Nitrogen, argon, etc.	19.0	16.8	15.0	13.5	12.3	11.3	10.4
Sum of above	29.2	25.7	22.8	20.5	18.7	17.1	15.7
% of oxygen in dissolved air (by vol.)	34.9%	34.7	34.5	34.2	34.0	33.8	33.6

GASES IN WATER

S indicates the number of c.c.s. of gas measured at 0° and 760 mms. which dissolve in 1 c.c. of water at the temperature stated, and when the pressure of the gas plus that of the water-vapour is 760 mms.

A indicates the same, except that the gas itself is at the uniform pressure of 760 mms. when in equilibrium with the water. (For other values, see p. 109.)

Gas.	0° C.	10°	15°	20°	30°	40°	50°	60°
	c.c.s.							
Ammonia, A	1300	910	802	710	595/28°	—	—	—
Argon, A058	.045	.040	.037	.030	.027	—	—
Carbon dioxide, A	1.713	1.194	1.019	.878	.66	.53	.44	.36
Carbon monoxide, A035	.028	.025	.023	.020	.018	.016	.015
Chlorine, S	—	3.09	2.63	2.26	1.77	1.41	1.20	1.0
Helium, A0150	.0144	.0139	.0138	.0138	.0139	.0140	—
Hydrogen, A0215	.0198	.0190	.0184	—	—	—	—
Hydrochloric acid, S	506	474	458	442	411	386	362	339
Nitrogen, A0239	.0196	.0179	.0164	.0138	.0118	.0106	.0100
Nitrous oxide, A	1.05/5°	.88	.74	.63	—	—	—	—
Nitric oxide, A074	.057	.051	.047	.040	.035	.031	.029
Oxygen, A049	.038	.034	.031	.026	.023	.021	.019
Sulphuretted hydrogen, A	4.68	3.52	3.05	2.67	—	—	—	—
Sulphur dioxide, S	79.8	56.6	47.3	39.4	27.2	18.8	—	—

Ne, .0147/20°; Kr, .0670 - .0788/20°; Xe, .1109/20° - Antropoff, 1910.

MUTUAL SOLUBILITIES OF LIQUIDS

The data for the uppermost layer of the two solutions in equilibrium are given in the first line in each case. The pressure in some cases exceeds one atmosphere. Numbers are grams per 100 grams of solution. (From data in Seidell's "Solubilities.")

Liquids.	0°C.	10°	20°	30°	40°	50°	60°	70°	80°	00			
{Water in ether; ethereal layer	1.0	1.1	1.2	1.3	1.5	1.7	1.8	2.0	2.2	—			
{Ether in water; aqueous layer	12	8.7	6.5	5.1	4.5	4.1	3.7	3.2	2.8	—			
{Aniline (C ₆ H ₅ NH ₂) in water; aqueous layer	—	—	3.2	—	3.5	—	3.8	—	4.5	6			
{Aniline in water; aniline layer	—	—	95.5	—	95	—	95	—	93	92			
{Phenol (C ₆ H ₅ OH) in water; aqueous layer	—	7.5	8.3	8.8	9.6	12	17	33.4	{ at crit. temp. 68° 3				
{Phenol in water; phenol layer	—	75	72	70	67	63	55	33.4					
{Triethylamine in water; amine layer	51.9	at	72	97	96	96	96	{ at crit. temp.					
{Triethylamine [N(C ₂ H ₅) ₃] in aqueous layer	51.9	18° 6	14.2	5.8	3.6	2.9	2.2						
{CS ₂ in methyl alcohol; alcoholic layer	—	45	51	58	80.5	{ at crit. temp.							
{CS ₂ in CH ₃ OH; carbon bisulphide layer	—	98	97	96	80.5						40° 5		

SOLUBILITIES OF SOLIDS IN WATER

s = number of grams of **anhydrous** substance which when dissolved in 100 grams of **water** make a saturated solution at the temperature stated.

p = no. of grams of anhydrous substance per 100 grams of saturated **solution**.

The formula given is that of the solid phase which is in equilibrium with the solution. (See Seidell's "Solubilities," New York, 1907, where the most complete and accurate data will be found for solubilities.) For other solutions, see p. 109.

Substance.		0° C.	10°	15°	20°	40°	60°	80°	100°
Am. chloride, NH_4Cl	s	29.4	33.3	35.2	37.2	45.8	55.2	65.6	77.3
Barium chloride, $\text{BaCl}_2 \cdot 2\text{H}_2\text{O}$	s	31.6	33.3	34.4	35.7	40.7	46.4	52.4	58.8
Barium hydrate, $\text{Ba}(\text{OH})_2 \cdot 8\text{H}_2\text{O}$	s	1.67	2.48	3.23	3.89	8.22	20.9	101.4	—
Bromine (<i>liquid</i>), Br.	s	4.22	3.4	3.25	3.20	—	—	—	—
Cadmium sulphate, $\text{CdSO}_4 \cdot 8/3\text{H}_2\text{O}$	s	76.5	76.0	76.3	76.6	78.5	83.7	69.7*	60.77*
Ca. hydrate, $\text{Ca}(\text{OH})_2$	s	185	176	170	165	141	116	99.4	97.7
Copper sulphate, $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$	s	14.3	17.4	18.8	20.7	28.5	40.0	55.0	75.0
Li. carbonate, Li_2CO_3	s	1.54	1.43	1.38	1.33	1.17	1.01	.850	.720
Merc. chloride, HgCl_2	p	3.50	4.50	5.00	5.40	9.30	14.0	23.1	38.0
Potass. chloride, KCl	s	27.6	31.0	32.4	34.0	40.0	45.5	51.1	56.7
Potass. bromide, KBr	s	53.5	59.5	62.5	65.2	75.5	85.5	95.0	104
Potassium iodide, KI	s	127.5	136	140	144	160	176	192	208
Potassium hydrate, $\text{KOH} \cdot 2\text{H}_2\text{O}$	s	97.0	103	107	112	138§	—	—	178§
Potass. nitrate, KNO_3	s	13.3	20.9	25.8	32	64	110	169	246
Silv. nitrate, AgNO_3	s	122	170	196	222	376	525	669	952
Sodium carbonate, $\text{Na}_2\text{CO}_3 \cdot 10\text{H}_2\text{O}$	s	7.0	12.5	16.4	21.5	46.1	46.0	45.8	45.5
Sod. chloride, NaCl	s	35.7	35.8	35.9	36.0	36.6	37	38	39.0
Sodium sulphate, $\text{Na}_2\text{SO}_4 \cdot 10\text{H}_2\text{O}$	s	5.0	9.0	13.4	19.4	49†	45†	44†	42†
Strontium chloride, $\text{SrCl}_2 \cdot 6\text{H}_2\text{O}$	s	43	48	50	53	65	82	91‡	101‡
Succinic acid, $(\text{CH}_2)_2(\text{COOH})_2$	s	2.80	4.50	5.7	6.9	16.2	35.8	70.8	125
Sugar (Cane), $\text{C}_{12}\text{H}_{22}\text{O}_{11}$	s	179	190	197	204	238	287	362	487

* Solid phase becomes $\text{CdSO}_4 \cdot \text{H}_2\text{O}$ at 74°.

† Becomes Na_2SO_4 at 32° 38.

‡ Becomes $\text{SrCl}_2 \cdot 2\text{H}_2\text{O}$ at 70°. § Becomes $\text{KOH} \cdot 3/2\text{H}_2\text{O}$ at 32° 5 and $\text{KOH} \cdot \text{H}_2\text{O}$ at 50°.

|| Becomes $\text{Na}_2\text{CO}_3 \cdot \text{H}_2\text{O}$ at 35°.

COMPOSITION OF DRY ATMOSPHERIC AIR

(Ramsay, *Proc. Roy. Soc.*, 1908 ; G. Claude, *Compt. Rend.*, 1909.)

	N_2	O_2	A	CO_2	Kr	Xe	Ne	He
By weight .	75.5	23.2	1.3	.046 to .4	.028	.005	.0386	.0156
By volume .	78.05	21.0*	.95	.03 to .3	—	—	.02123	.0340

* 20.91 according to Kreusler.

MINERALS

MOHS' SCALE OF MINERAL HARDNESS

The numbers are not quantitative, but merely indicate the sequence of hardness.

Hardness.	Mineral.	Hardness.	Mineral.	Hardness.	Mineral.
1	Talc	5	Apatite	9	Corundum
2	Rock salt	6	Felspar	10	Diamond
3	Calcspars	7	Quartz	<i>c. 2.5</i>	Finger-nail
4	Fluor spar	8	Topaz	<i>c. 6.5</i>	Penknife

COMPOSITION, DENSITY, AND HARDNESS OF SOME MINERALS

See Dana's "System of Mineralogy" and Appendices, 1892, 1899, and 1909. Radioactive minerals are indicated thus *; see Szilard, *Le Radium*, August, 1909.

Name and Formula.	Density.	Hardness.	Name and Formula.	Density.	Hardness.
Albite, $\text{Na}_2\text{Al}_2\text{Si}_6\text{O}_{16}$. . .	<i>c. 2.6</i>	6-7	Mica (common, Muscovite), $\text{K}_2\text{O} \cdot 3\text{Al}_2\text{O}_3 \cdot 6\text{SiO}_2 \cdot 2\text{H}_2\text{O}$	2.7-3.1	2-2.5
Amber (fossil resin) . . .	1.08	2-2.5	Mica (Biotite, Magnesia mica)	2.7-3.1	2.5-3
Anhydrite, CaSO_4 . . .	2.8-2.9	3-3.5	Monazite,* $(\text{CeLaDi})\text{PO}_4$ (1-16% Th)	5	5.2
Anorthite, $\text{Ca}_2\text{Al}_2\text{Si}_2\text{O}_{16}$. .	<i>c. 2.7</i>	6-7	Nepheline, $\text{Na}_4\text{K}_6\text{Al}_3\text{Si}_9\text{O}_{36}$	2.5-2.6	5.5-6
Apatite, $\text{Ca}_5(\text{Cl},\text{F},\text{OH})(\text{PO}_4)_3$	2.9-3.2	5	Olivine, $\text{Mg}_2\text{Fe}_2\text{SiO}_4$. . .	3.3-3.5	6-7
Aragonite, CaCO_3 . . .	2.93	3.5-4	Orthoclase, $\text{K}_2\text{Al}_2\text{Si}_6\text{O}_{16}$. . .	2.4-2.6	6
Augite, Mg, Fe, Ca, Al silicate	3.2-3.5	5-6	Pitchblende,* U_3O_8 with oxides of Pb, and Ca, Fe, Bi, Mn, Mg, Cu, Si, Al, etc. (25-80% U; 1-6% Th)	6.4 (mas- sive) 9.7 (cryst.)	5.5
Barytes, Heavy spar, BaSO_4	4.5	3-3.5	Pyrites (iron), FeS_2 . . .	4.8-5.1	6-6.5
Beryl, $\text{Be}_3\text{Al}_2\text{Si}_6\text{O}_{18}$. . .	2.6-2.7	7-8	" (copper), CuFeS_2 . . .	4.1-4.3	3.5-4
Bröggerite,* a pitch- blende which contains thorium	(56-68% U)	(2-8% Th)	Pyrolusite, MnO_2 . . .	4.8-5	2-5.5
Calcite, Calcspars, Iceland spar, CaCO_3	2.6-2.7	<i>c. 3</i>	Quartz, SiO_2 . . .	2.5-2.8	7
Carnallite, $\text{KCl} \cdot \text{MgCl}_2 \cdot 6\text{H}_2\text{O}$	1.6	1	Rock salt, NaCl . . .	2.1-2.2	2-2.5
Carnotite,* $\text{K}_2\text{O}(\text{UO}_2)_2\text{V}_2\text{O}_6 \cdot 3\text{H}_2\text{O}$	(<i>c. 55%</i> U)	(yel- low)	Rutile, TiO_2 . . .	4.2-4.3	6-6.5
Celestine, SrSO_4 . . .	3.9	3-3.5	Selenite—cryst. gypsum	—	—
Cerussite, PbCO_3 . . .	6.4	3-3.5	Serpentine, $\text{H}_2\text{Mg}_3\text{Si}_2\text{O}_9$	<i>c. 2.6</i>	3-4
Chalcocite,* $\text{Cu}(\text{UO}_2)(\text{PO}_4)_2 \cdot 8\text{H}_2\text{O}$	3.4-3.6 (48% U)	2-2.5	Spinel, MgOAl_2O_3 . . .	3.5-3.6	8
Cléveite*—pitchblende which contains Th & Y	(<i>c. 60%</i> U)	(<i>c. 4%</i> Th)	Sylvine, KCl . . .	1.9-2	2
Corundum, Al_2O_3 . . .	3.9-4.2	9	Talc, $\text{H}_2\text{Mg}_3\text{Si}_4\text{O}_{12}$. . .	2.5-2.8	1
Dolomite, CaMgC_2O_6 . . .	2.8-2.9	3.5-4	Thorianite,* Th, U ox- ides, etc.; (4-10% U; <i>c. 60%</i> Th) contains He	8-9.7	7 (black cubes)
Felspar, $\text{Al}_2\text{K}_2\text{Si}_6\text{O}_{16}$. . .	2.4-2.6	6	Thorite,* ThSiO_4 (1-9% U; 40-60% Th)	4.6	4.6 (tetra- gonal)
Flint; agate, SiO_2 . . .	2.6	<i>c. 6</i>	Tourmaline, hydrated sil- icate and borate of Al, Na with Li or Fe or Mg	2.9-3.3	7-7.5
Fluorspar, Fluorite, CaF_2	3-3.3	4	Trögerite,* $(\text{UO}_2)_3\text{As}_2\text{O}_5 \cdot 12\text{H}_2\text{O}$	(53% U)	(yel- low)
Galena, PbS . . .	7.4-7.6	2-3	Uraninite*—crystalline pitchblende (<i>q.v.</i>)	(Black)	octahe- dra
Gummite,* $\text{Pb}, \text{Ca}, \text{U}, \text{silic}$ ate(50-65% U)	2.3	6.5-7	Uranite lime,* $\text{CaO}(\text{UO}_2)_2(\text{PO}_4)_2 \cdot 8\text{H}_2\text{O}$ (50% U)	3-3.2	2-2.5
Gypsum, $\text{CaSO}_4 \cdot 2\text{H}_2\text{O}$. . .	1.5-2	1.5-2	Willemite, Zn_2SiO_4 . . .	4	5
Hæmatite, Fe_2O_3 . . .	4.5-5.3	5.5-6.5	Wolfram, (Fe, Mn) WO_4 . .	7.1-7.9	5-5.5
Hornblende, Ca, Mg, Fe, Na, Al, silicate	2.9-3.4	5-6	Wollastonite, CaSiO_3 . . .	2.7-2.9	4.5-5
Kainite, $\text{MgSO}_4 \cdot \text{KCl} \cdot 3\text{H}_2\text{O}$	2.1	—	Zeunerite,* Cu, U arse- nate (<i>c. 50%</i> U)	(<i>c. 50%</i> U)	(tetra- gonal)
Kaolin, $\text{H}_4\text{Al}_2\text{Si}_2\text{O}_9$. . .	2.5	1	Zircon,* ZrSiO_4 . . .	4.7	7.5
Kieserite, $\text{MgSO}_4 \cdot \text{H}_2\text{O}$. . .	2.55	3	Zincblende, ZnS . . .	3.9-4.2	3.5-4
Lepidolite (Lithia mica), $(\text{F}, \text{OH})_2(\text{Li}, \text{K}, \text{Na})_2\text{Al}_2\text{Si}_2\text{O}_9$	2.8-3	2.5-4			
Limestone, CaCO_3 . . .	2.5-2.8	—			
Magnesite, MgCO_3 . . .	<i>c. 3</i>	3.5-4.5			
Magnetite, Fe_3O_4 . . .	4.9-5.2	5.5-6.5			
Meerschaum, $2\text{MgO} \cdot 3\text{SiO}_2 \cdot 2\text{H}_2\text{O}$	<i>c. 2.6</i>	2-2.5			

FACTORS FOR GRAVIMETRIC ANALYSIS

Calculated with atomic weights for 1911 (p. 1).

Example.—1 gram Al_2O_3 is chemically equivalent to .5303 gram Al, or 1 gram Al is equivalent to $1/.5303 \text{ Al}_2\text{O}_3$. A table of reciprocals is given on p. 136. (See Van Nostrand's "Chemical Annual," London.)

1 part by weight of	is equivalent (by weight) to	1 part by weight of	is equivalent (by weight) to
Aluminium.		Calcium (<i>contd.</i>)—	
Al_2O_35303 Al	$\text{Ca}_3(\text{PO}_4)_2$5422 CaO
"	$3.350 \text{ Al}_2(\text{SO}_4)_3$	$\text{Mg}_3\text{P}_2\text{O}_7$	$1.3935 \text{ Ca}_3(\text{PO}_4)_2$
Ammonium.		P_2O_5	$2.1844 \text{ Ca}_3(\text{PO}_4)_2$
N	1.216 NH_3	Carbon.	
"	1.288 NH_4	CO_2	4.4860 BaCO_3
"	$3.819 \text{ NH}_4\text{Cl}$	"	2.2748 CaCO_3
NH_3	$2.058 \text{ NH}_4\text{OH}$	Chlorine.	
Antimony.		AgCl2474 Cl
Sb	$1.1997 \text{ Sb}_2\text{O}_3$	NaCl6066 Cl
"	$1.3328 \text{ Sb}_2\text{O}_5$	Chromium.	
Sb_2O_3	$1.1109 \text{ Sb}_2\text{O}_5$	Cr_2O_36846 Cr
Sb_2O_47897 Sb	"	1.3154 CrO_3
"	$.9474 \text{ Sb}_2\text{O}_3$	Cobalt.	
"	$1.0526 \text{ Sb}_2\text{O}_5$	Co	1.2713 CoO
Arsenic.		Co_3O_47343 Co
As_2O_37575 As	"9336 CoO
"	$1.1617 \text{ As}_2\text{O}_5$	Co(NO_2) $_3$. (KNO_2) $_3$.1306 Co
As_2O_56521 As	"1661 CoO
$\text{MgNH}_4\text{AsO}_4 \cdot \frac{1}{2} \text{H}_2\text{O}$.3938 As	(CoSO_4) $_2$. (K_2SO_4) $_3$.1416 Co
"5199 As_2O_3	Copper.	
"6040 As_2O_5	Cu	1.2517 CuO
$\text{Mg}_3\text{As}_2\text{O}_7$4827 As	Fluorine.	
"6373 As_2O_3	CaF_24866 F
"7403 As_2O_5	Glucinum. See	
Barium.		Beryllium.	
BaCO_36960 Ba	Gold.	
"7771 BaO	Au	1.5395 AuCl_3
BaSO_45885 Ba	Hydrogen.	
"6570 BaO	H_2O1119 H
"7255 BaO_2	Iodine.	
Beryllium.		AgI5405 I
BeO3626 Be	Iron.	
Bismuth.		Fe	1.2865 FeO
Bi	$1.1154 \text{ Bi}_2\text{O}_3$	"	$1.4297 \text{ Fe}_2\text{O}_3$
Bi_2O_38966 Bi	"	$7.0218 \text{ FeSO}_4 \cdot (\text{NH}_4)_2\text{SO}_4 \cdot 6\text{H}_2\text{O}$
BiOCl8017 Bi	FeO7773 Fe
"8942 Bi_2O_3	"	$1.1113 \text{ Fe}_2\text{O}_3$
Boron.		Fe_2O_3	1.4508 FeCO_3
B_2O_33143 B	"9666 Fe_2O_4
"	$2.7297 \text{ Na}_3\text{B}_4\text{O}_7 \cdot 10\text{H}_2\text{O}$	CO_2	1.6330 FeO
Bromine.		"	2.6330 FeCO_3
AgBr4256 Br	Lead.	
Cadmium.		Pb	1.0773 PbO
CdO8754 Cd	PbSO_46831 Pb
Cæsium.		"7358 PbO
Cs	$1.060 \text{ Cs}_2\text{O}$	"7887 PbO_2
Cs_2PtCl_63945 Cs	"7536 Pb_3O_4
"4184 Cs_2O	Lithium.	
Calcium.		Li_2CO_31879 Li
Ca	1.399 CaO	"4044 Li_2O
CaCO_34005 Ca	Li_3PO_41797 Li
"5604 CaO	"3868 Li_2O
CO_2	2.275 CaCO_3		

GRAVIMETRIC FACTORS

FACTORS FOR GRAVIMETRIC ANALYSIS (contd.)

1 part by weight of	is equivalent (by weight) to	1 part by weight of	is equivalent (by weight) to
Magnesium.		Potassium (contd.)	
MgO	·6032 Mg	K ₂ SO ₄	1·1604 KNO ₃
Mg ₂ P ₂ O ₇	·2184 Mg	K ₂ PtCl ₆	·1609 K
"	·3621 MgO	Rubidium.	
Manganese.		Rb ₂ PtCl ₆	·2953 Rb
MnO	1·1113 Mn ₂ O ₃	Silicon.	
Mn ₂ O ₃	·7203 Mn	SiO ₂	·4693 Si
"	·9307 MnO	Silver.	
"	1·0350 Mn ₂ O ₃	AgCl	·7526 Ag
"	1·1399 MnO ₂	AgBr	·5744 Ag
Mercury.		AgI	·4595 Ag
Hg	1·1603 HgS	Sodium.	
HgS	·8963 Hg ₂ O	AgCl	·4078 NaCl
"	·9308 HgO	NaHCO ₃	·3691 Na ₂ O
Nickel.		Na ₂ SO ₄	·3238 Na
Ni	1·2727 NiO	"	·4364 Na ₂ O
Nitrogen.		N ₂ O ₅	1·5740 NaNO ₃
N	3·8551 N ₂ O ₅	Strontium.	
Phosphorus.		SrCO ₃	·7019 SrO
P ₂ O ₅	·4362 P	SrSO ₄	·5641 SrO
Mg ₂ P ₂ O ₇	·2787 P	Sulphur.	
"	·8534 PO ₄	BaSO ₄	·1460 H ₂ S
"	·6378 P ₂ O ₅	"	·1374 S
Platinum.		"	·2744 SO ₂
K ₂ PtCl ₆	·4015 Pt	"	·3429 SO ₃
"	·6933 PtCl ₄	"	·4115 SO ₄
Potassium.		Tin.	
AgCl	·5202 KCl	SnO ₂	·7881 Sn
AgBr	·6338 KBr	Uranium.	
AgI	·7071 KI	U ₃ O ₈	·8482 U
AgCN	·4863 KCN	"	·9620 UO ₂
KCl	·5244 K	UO ₂	·8817 U
KBr	·3285 K	Zinc.	
KOH	1·2316 K ₂ CO ₃	Zn	1·2448 ZnO
"	·8395 K ₂ O	ZnO	·8033 Zn
K ₂ SO ₄	·5403 K ₂ O		

SOME BOILING-POINT MIXTURES

Boiling-points under 760 mms. of mercury. Percentage compositions by weight. A large number of minimum boiling-point mixtures are known.

(Sidney Young, "Fractional Distillation," 1903.)

	Mixture.		Boiling Points.			% of A in mixt.	Ob- server.
	A.	B.	A.	B.	Mixt.		
Maximum boiling- point mixtures.	Water	Nitric acid	100° C.	86°	125°	32%	Roscoe
	"	Hydrochloric acid	100	c. - 80	110	80	"
	"	Formic acid	100	100·8	107	23	"
	Me. ether	Hydrochloric acid	- 23·6	c. - 80	- 2	61	Friedel
Minimum boiling- point mixtures.	Water	Ethyl alcohol	100	78·3	78·1	4·4	Y. & F.
	Pyridine	Water	117	100	92·5	59	G. & C.
	Benzene	Methyl alcohol	80·2	64·7	58·3	60	Y. & F.
	Me. alcohol	Acetone	64·7	- 56·5	55·9	13·5	Pettit

G. & C., Goldschmidt and Constan ; Y. & F., Young and Fortey.

THE EXPONENTIAL e^{-x} $e = 2.71828$. To derive e^x use reciprocals on p. 136. $e^{-.69315} = .5$.(Based on Newman, *Trans. Camb. Phil. Soc.*, 13, 1883.)

For values of x from .0000 to .0999.											Subtract Differences.									
x	0	.001	.002	.003	.004	.005	.006	.007	.008	.009	.0001	2	3	4	5	6	7	8	9	
.00	1.000	.9990	.9980	.9970	.9960	.9950	.9940	.9930	.9920	.9910	1	2	3	4	5	6	7	8	9	
.01	.9900	.9891	.9881	.9871	.9861	.9851	.9841	.9831	.9822	.9812	1	2	3	4	5	6	7	8	9	
.02	.9802	.9792	.9782	.9773	.9763	.9753	.9743	.9734	.9724	.9714	1	2	3	4	5	6	7	8	9	
.03	.9704	.9695	.9685	.9675	.9666	.9656	.9646	.9637	.9627	.9618	1	2	3	4	5	6	7	8	9	
.04	.9608	.9598	.9589	.9579	.9570	.9560	.9550	.9541	.9531	.9522	1	2	3	4	5	6	7	8	9	
.05	.9512	.9502	.9493	.9484	.9474	.9465	.9455	.9446	.9436	.9427	1	2	3	4	5	6	7	8	9	
.06	.9418	.9408	.9399	.9389	.9380	.9371	.9361	.9352	.9343	.9333	1	2	3	4	5	6	7	8	9	
.07	.9324	.9315	.9305	.9296	.9287	.9277	.9268	.9259	.9250	.9240	1	2	3	4	5	6	7	8	8	
.08	.9231	.9222	.9213	.9204	.9194	.9185	.9176	.9167	.9158	.9148	1	2	3	4	5	6	7	7	8	
.09	.9139	.9130	.9121	.9112	.9103	.9094	.9085	.9076	.9066	.9057	1	2	3	4	5	6	6	7	8	

For values of x from .100 to 2.999.											Subtract Differences.									
x	0	.01	.02	.03	.04	.05	.06	.07	.08	.09	.001	2	3	4	5	6	7	8	9	
.1	.9048	.8958	.8866	.8781	.8694	.8607	.8521	.8437	.8353	.8270	9	17	26	34	43	52	60	69	77	
.2	.8187	.8106	.8025	.7945	.7866	.7788	.7711	.7634	.7558	.7483	8	16	23	31	39	47	55	62	70	
.3	.7408	.7334	.7261	.7189	.7118	.7047	.6977	.6907	.6839	.6771	7	14	21	28	35	42	49	56	63	
.4	.6703	.6637	.6570	.6505	.6440	.6376	.6313	.6250	.6188	.6126	6	13	19	26	32	38	45	51	57	
.5	.6065	.6005	.5945	.5886	.5827	.5769	.5712	.5655	.5599	.5543	6	12	17	23	29	35	40	46	52	
.6	.5488	.5434	.5379	.5326	.5273	.5220	.5169	.5117	.5066	.5016	5	10	16	21	26	31	37	42	47	
.7	.4966	.4916	.4868	.4819	.4771	.4724	.4677	.4630	.4584	.4538	5	9	14	19	24	28	33	38	43	
.8	.4493	.4449	.4404	.4360	.4317	.4274	.4232	.4190	.4148	.4107	4	9	13	17	21	26	30	34	38	
.9	.4066	.4025	.3985	.3946	.3906	.3867	.3829	.3791	.3753	.3716	4	8	12	15	19	23	27	31	35	
1.0	.3679	.3642	.3606	.3570	.3535	.3499	.3465	.3430	.3396	.3362	4	7	11	14	18	21	25	28	32	
1.1	.3329	.3296	.3263	.3230	.3198	.3166	.3135	.3104	.3073	.3042	3	6	9	13	16	19	22	25	29	
1.2	.3012	.2982	.2952	.2923	.2894	.2865	.2837	.2808	.2780	.2753	3	6	9	11	14	17	20	23	26	
1.3	.2725	.2698	.2671	.2645	.2618	.2592	.2567	.2541	.2516	.2491	3	5	8	10	13	16	18	21	23	
1.4	.2466	.2441	.2417	.2393	.2369	.2346	.2322	.2299	.2276	.2254	2	5	7	9	12	14	16	19	21	
1.5	.2231	.2209	.2187	.2165	.2144	.2122	.2101	.2080	.2060	.2039	2	4	6	8	11	13	15	17	19	
1.6	.2019	.1999	.1979	.1959	.1940	.1920	.1901	.1882	.1864	.1845	2	4	6	8	10	12	13	15	17	
1.7	.1827	.1809	.1791	.1773	.1755	.1738	.1720	.1703	.1686	.1670	2	3	5	7	9	10	12	14	16	
1.8	.1653	.1637	.1620	.1604	.1588	.1572	.1557	.1541	.1526	.1511	2	3	5	6	8	9	11	13	14	
1.9	.1496	.1481	.1466	.1451	.1437	.1423	.1409	.1395	.1381	.1367	1	3	4	6	7	9	10	11	13	
2.0	.1353	.1340	.1327	.1313	.1300	.1287	.1275	.1262	.1249	.1237	1	3	4	5	6	8	9	10	12	
2.1	.1225	.1212	.1200	.1188	.1177	.1165	.1153	.1142	.1130	.1119	1	2	4	5	6	7	8	9	11	
2.2	.1108	.1097	.1086	.1075	.1065	.1054	.1044	.1033	.1023	.1013	1	2	3	4	5	6	7	8	9	
2.3	.1003	.0993	.0983	.0973	.0963	.0954	.0944	.0935	.0926	.0916	1	2	3	4	5	6	7	8	9	
2.4	.0907	.0898	.0889	.0880	.0872	.0863	.0854	.0846	.0837	.0829	1	2	3	4	5	6	7	8	9	
2.5	.0821	.0813	.0805	.0797	.0789	.0781	.0773	.0765	.0758	.0750	1	2	3	4	5	5	6	7	8	
2.6	.0743	.0735	.0728	.0721	.0714	.0707	.0699	.0693	.0686	.0679	1	1	2	3	4	4	5	6	6	
2.7	.0672	.0665	.0659	.0652	.0646	.0639	.0633	.0627	.0620	.0614	1	1	2	3	3	4	4	5	6	
2.8	.0608	.0602	.0596	.0590	.0584	.0578	.0573	.0567	.0561	.0556	1	1	2	2	3	3	4	4	5	
2.9	.0550	.0545	.0539	.0534	.0529	.0523	.0518	.0513	.0508	.0503	1	1	2	2	3	3	4	4	5	

For values of x from 3.0 to 8.9.											Subtract Differences.									
x	0	.1	.2	.3	.4	.5	.6	.7	.8	.9	Mean differences no longer sufficiently accurate.									
3	.0498	.0450	.0408	.0368	.0334	.0302	.0273	.0247	.0224	.0202										
4	.0183	.0166	.0150	.0136	.0123	.0111	.0101	.0091	.0082	.0074										
5	.0067	.0061	.0055	.0050	.0045	.0041	.0037	.0033	.0030	.0027										
6	.0025	.0022	.0020	.0018	.0017	.0015	.0014	.0012	.0011	.0010										
7	.0009	.0008	.0007	.0007	.0006	.0006	.0005	.0005	.0004	.0004										
8	.0003	.0003	.0003	.0002	.0002	.0002	.0002	.0002	.0002	.0001										

FOUR-FIGURE LOGARITHMS

	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9
10	0000	0043	0086	0128	0170	0212	0253	0294	0334	0374	4	9	13	17	21	25	30	34	38
11	0414	0453	0492	0531	0569	0607	0645	0682	0719	0755	4	8	12	15	19	23	27	31	35
12	0792	0828	0864	0899	0934	0969	1004	1038	1072	1106	4	7	11	14	18	21	25	28	32
13	1139	1173	1206	1239	1271	1303	1335	1367	1399	1430	3	7	10	13	16	20	23	26	30
14	1461	1492	1523	1553	1584	1614	1644	1673	1703	1732	3	6	9	12	15	18	21	24	27
15	1761	1790	1818	1847	1875	1903	1931	1959	1987	2014	3	6	8	11	14	17	20	23	26
16	2041	2068	2095	2122	2148	2175	2201	2227	2253	2279	3	5	8	11	13	16	19	21	24
17	2304	2330	2355	2380	2405	2430	2455	2480	2504	2529	3	5	8	10	13	16	18	21	23
18	2553	2577	2601	2625	2648	2672	2695	2718	2742	2765	2	5	7	10	12	15	17	20	22
19	2788	2810	2833	2856	2878	2900	2923	2945	2967	2989	2	5	7	9	11	14	16	18	20
20	3010	3032	3054	3075	3096	3118	3139	3160	3181	3201	2	4	6	8	11	13	15	17	19
21	3222	3243	3263	3284	3304	3324	3345	3365	3385	3404	2	4	6	8	10	12	14	16	18
22	3424	3444	3464	3483	3502	3522	3541	3560	3579	3598	2	4	6	8	10	12	14	15	17
23	3617	3636	3655	3674	3692	3711	3729	3747	3766	3784	2	4	6	7	9	11	13	15	17
24	3802	3820	3838	3856	3874	3892	3909	3927	3945	3962	2	4	5	7	9	11	12	14	16
25	3979	3997	4014	4031	4048	4065	4082	4099	4116	4133	2	3	5	7	9	10	12	14	15
26	4150	4166	4183	4200	4216	4232	4249	4265	4281	4298	2	3	5	7	8	10	11	13	15
27	4314	4330	4346	4362	4378	4393	4409	4425	4440	4456	2	3	5	6	8	9	11	12	14
28	4472	4487	4502	4518	4533	4548	4564	4579	4594	4609	2	3	5	6	8	9	11	13	14
29	4624	4639	4654	4669	4683	4698	4713	4728	4742	4757	1	3	4	6	7	9	10	12	13
30	4771	4786	4800	4814	4829	4843	4857	4871	4886	4900	1	3	4	6	7	9	10	11	13
31	4914	4928	4942	4955	4969	4983	4997	5011	5024	5038	1	3	4	6	7	8	10	11	12
32	5051	5065	5079	5092	5105	5119	5132	5145	5159	5172	1	3	4	5	7	8	9	11	12
33	5185	5198	5211	5224	5237	5250	5263	5276	5289	5302	1	3	4	5	6	8	9	10	12
34	5315	5328	5340	5353	5366	5378	5391	5403	5416	5428	1	3	4	5	6	8	9	10	11
35	5441	5453	5465	5478	5490	5502	5514	5527	5539	5551	1	2	4	5	6	7	9	10	11
36	5563	5575	5587	5599	5611	5623	5635	5647	5658	5670	1	2	4	5	6	7	8	10	11
37	5682	5694	5705	5717	5729	5740	5752	5763	5775	5786	1	2	3	5	6	7	8	9	10
38	5798	5809	5821	5832	5843	5855	5866	5877	5888	5899	1	2	3	5	6	7	8	9	10
39	5911	5922	5933	5944	5955	5966	5977	5988	5999	6010	1	2	3	4	5	7	8	9	10
40	6021	6031	6042	6053	6064	6075	6085	6096	6107	6117	1	2	3	4	5	6	8	9	10
41	6128	6138	6149	6160	6170	6180	6191	6201	6212	6222	1	2	3	4	5	6	7	8	9
42	6232	6243	6253	6263	6274	6284	6294	6304	6314	6325	1	2	3	4	5	6	7	8	9
43	6335	6345	6355	6365	6375	6385	6395	6405	6415	6425	1	2	3	4	5	6	7	8	9
44	6435	6444	6454	6464	6474	6484	6493	6503	6513	6522	1	2	3	4	5	6	7	8	9
45	6532	6542	6551	6561	6571	6580	6590	6599	6609	6618	1	2	3	4	5	6	7	8	9
46	6628	6637	6646	6656	6665	6675	6684	6693	6702	6712	1	2	3	4	5	6	7	7	8
47	6721	6730	6739	6749	6758	6767	6776	6785	6794	6803	1	2	3	4	5	5	6	7	8
48	6812	6821	6830	6839	6848	6857	6866	6875	6884	6893	1	2	3	4	4	5	6	7	8
49	6902	6911	6920	6928	6937	6946	6955	6964	6972	6981	1	2	3	4	4	5	6	7	8
	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9

FOUR-FIGURE LOGARITHMS

	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9
50	6990	6998	7007	7016	7024	7033	7042	7050	7059	7067	1	2	3	3	4	5	6	7	8
51	7076	7084	7093	7101	7110	7118	7126	7135	7143	7152	1	2	3	3	4	5	6	7	8
52	7160	7168	7177	7185	7193	7202	7210	7218	7226	7235	1	2	2	3	4	5	6	7	7
53	7243	7251	7259	7267	7275	7284	7292	7300	7308	7316	1	2	2	3	4	5	6	6	7
54	7324	7332	7340	7348	7356	7364	7372	7380	7388	7396	1	2	2	3	4	5	6	6	7
55	7404	7412	7419	7427	7435	7443	7451	7459	7466	7474	1	2	2	3	4	5	5	6	7
56	7482	7490	7497	7505	7513	7520	7528	7536	7543	7551	1	2	2	3	4	5	5	6	7
57	7559	7566	7574	7582	7589	7597	7604	7612	7619	7627	1	2	2	3	4	5	5	6	7
58	7634	7642	7649	7657	7664	7672	7679	7686	7694	7701	1	1	2	3	4	4	5	6	7
59	7709	7716	7723	7731	7738	7745	7752	7760	7767	7774	1	1	2	3	4	4	5	6	7
60	7782	7789	7796	7803	7810	7818	7825	7832	7839	7846	1	1	2	3	4	4	5	6	6
61	7853	7860	7868	7875	7882	7889	7896	7903	7910	7917	1	1	2	3	4	4	5	6	6
62	7924	7931	7938	7945	7952	7959	7966	7973	7980	7987	1	1	2	3	3	4	5	6	6
63	7993	8000	8007	8014	8021	8028	8035	8041	8048	8055	1	1	2	3	3	4	5	5	6
64	8062	8069	8075	8082	8089	8096	8102	8109	8116	8122	1	1	2	3	3	4	5	5	6
65	8129	8136	8142	8149	8156	8162	8169	8176	8182	8189	1	1	2	3	3	4	5	5	6
66	8195	8202	8209	8215	8222	8228	8235	8241	8248	8254	1	1	2	3	3	4	5	5	6
67	8261	8267	8274	8280	8287	8293	8299	8306	8312	8319	1	1	2	3	3	4	5	5	6
68	8325	8331	8338	8344	8351	8357	8363	8370	8376	8382	1	1	2	3	3	4	4	5	6
69	8388	8395	8401	8407	8414	8420	8426	8432	8439	8445	1	1	2	3	3	4	4	5	6
70	8451	8457	8463	8470	8476	8482	8488	8494	8500	8506	1	1	2	2	3	4	4	5	6
71	8513	8519	8525	8531	8537	8543	8549	8555	8561	8567	1	1	2	2	3	4	4	5	5
72	8573	8579	8585	8591	8597	8603	8609	8615	8621	8627	1	1	2	2	3	4	4	5	5
73	8633	8639	8645	8651	8657	8663	8669	8675	8681	8686	1	1	2	2	3	4	4	5	5
74	8692	8698	8704	8710	8716	8722	8727	8733	8739	8745	1	1	2	2	3	4	4	5	5
75	8751	8756	8762	8768	8774	8779	8785	8791	8797	8802	1	1	2	2	3	3	4	5	5
76	8808	8814	8820	8825	8831	8837	8842	8848	8854	8859	1	1	2	2	3	3	4	5	5
77	8865	8871	8876	8882	8887	8893	8899	8904	8910	8915	1	1	2	2	3	3	4	4	5
78	8921	8927	8932	8938	8943	8949	8954	8960	8965	8971	1	1	2	2	3	3	4	4	5
79	8976	8982	8987	8993	8998	9004	9009	9015	9020	9025	1	1	2	2	3	3	4	4	5
80	9031	9036	9042	9047	9053	9058	9063	9069	9074	9079	1	1	2	2	3	3	4	4	5
81	9085	9090	9096	9101	9106	9112	9117	9122	9128	9133	1	1	2	2	3	3	4	4	5
82	9138	9143	9149	9154	9159	9165	9170	9175	9180	9186	1	1	2	2	3	3	4	4	5
83	9191	9196	9201	9206	9212	9217	9222	9227	9232	9238	1	1	2	2	3	3	4	4	5
84	9243	9248	9253	9258	9263	9269	9274	9279	9284	9289	1	1	2	2	3	3	4	4	5
85	9294	9299	9304	9309	9315	9320	9325	9330	9335	9340	1	1	2	2	3	3	4	4	5
86	9345	9350	9355	9360	9365	9370	9375	9380	9385	9390	1	1	2	2	3	3	4	4	5
87	9395	9400	9405	9410	9415	9420	9425	9430	9435	9440	0	1	1	2	2	3	3	4	4
88	9445	9450	9455	9460	9465	9469	9474	9479	9484	9489	0	1	1	2	2	3	3	4	4
89	9494	9499	9504	9509	9513	9518	9523	9528	9533	9538	0	1	1	2	2	3	3	4	4
90	9542	9547	9552	9557	9562	9566	9571	9576	9581	9586	0	1	1	2	2	3	3	4	4
91	9590	9595	9600	9605	9609	9614	9619	9624	9628	9633	0	1	1	2	2	3	3	4	4
92	9638	9643	9647	9652	9656	9661	9666	9671	9675	9680	0	1	1	2	2	3	3	4	4
93	9685	9689	9694	9699	9703	9708	9713	9717	9722	9727	0	1	1	2	2	3	3	4	4
94	9731	9736	9741	9745	9750	9754	9759	9763	9768	9773	0	1	1	2	2	3	3	4	4
95	9777	9782	9786	9791	9795	9800	9805	9809	9814	9818	0	1	1	2	2	3	3	4	4
96	9823	9827	9832	9836	9841	9845	9850	9854	9859	9863	0	1	1	2	2	3	3	4	4
97	9868	9872	9877	9881	9886	9890	9894	9899	9903	9908	0	1	1	2	2	3	3	4	4
98	9912	9917	9921	9926	9930	9934	9939	9943	9948	9952	0	1	1	2	2	3	3	4	4
99	9956	9961	9965	9969	9974	9978	9983	9987	9991	9996	0	1	1	2	2	3	3	3	4
	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9

ANTILOGARITHMS

	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9
00	1000	1002	1005	1007	1009	1012	1014	1016	1019	1021	0	0	1	1	1	1	2	2	2
01	1023	1026	1028	1030	1033	1035	1038	1040	1042	1045	0	0	1	1	1	1	2	2	2
02	1047	1050	1052	1054	1057	1059	1062	1064	1067	1069	0	0	1	1	1	1	2	2	2
03	1072	1074	1076	1079	1081	1084	1086	1089	1091	1094	0	0	1	1	1	1	2	2	2
04	1096	1099	1102	1104	1107	1109	1112	1114	1117	1119	0	1	1	1	1	1	2	2	2
05	1122	1125	1127	1130	1132	1135	1138	1140	1143	1146	0	1	1	1	1	1	2	2	2
06	1148	1151	1153	1156	1159	1161	1164	1167	1169	1172	0	1	1	1	1	1	2	2	2
07	1175	1178	1180	1183	1186	1189	1191	1194	1197	1199	0	1	1	1	1	1	2	2	2
08	1202	1205	1208	1211	1213	1216	1219	1222	1225	1227	0	1	1	1	1	1	2	2	3
09	1230	1233	1236	1239	1242	1245	1247	1250	1253	1256	0	1	1	1	1	1	2	2	3
10	1259	1262	1265	1268	1271	1274	1276	1279	1282	1285	0	1	1	1	1	1	2	2	3
11	1288	1291	1294	1297	1300	1303	1306	1309	1312	1315	0	1	1	1	1	1	2	2	3
12	1318	1321	1324	1327	1330	1334	1337	1340	1343	1346	0	1	1	1	1	1	2	2	3
13	1349	1352	1355	1358	1361	1365	1368	1371	1374	1377	0	1	1	1	1	1	2	2	3
14	1380	1384	1387	1390	1393	1396	1400	1403	1406	1409	0	1	1	1	1	1	2	2	3
15	1413	1416	1419	1422	1426	1429	1432	1435	1439	1442	0	1	1	1	1	1	2	2	3
16	1445	1449	1452	1455	1459	1462	1466	1469	1472	1476	0	1	1	1	1	1	2	2	3
17	1479	1483	1486	1489	1493	1496	1500	1503	1507	1510	0	1	1	1	1	1	2	2	3
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FIVE-FIGURE LOGARITHMS

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						02119	02531	02938	03342	03743	41	81	121	162	202	243	283	323	364
11	04139	04532	04922	05308	05690						39	77	116	155	193	232	270	309	348
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12	07918	08279	08636	08991	09342						36	71	106	142	177	213	248	284	319
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13	11394	11727	12057	12385	12710						33	66	98	131	164	197	230	262	295
						13033	13354	13672	13988	14301	32	63	95	126	158	190	221	253	284
14	14613	14922	15229	15534	15836						31	61	91	122	152	183	213	244	274
						16137	16435	16732	17026	17319	30	59	88	118	147	177	206	236	265
15	17609	17898	18184	18469	18752						29	57	85	114	142	171	199	228	256
						19033	19312	19590	19866	20140	28	55	83	110	138	166	193	221	248
16	20412	20683	20951	21219	21484						27	53	80	107	134	160	187	214	241
						21748	22011	22272	22531	22789	26	52	78	104	130	156	182	208	233
17	23045	23300	23553	23805	24055						25	50	76	101	126	151	176	201	227
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18	25527	25768	26007	26245	26482						24	48	71	95	119	143	167	190	214
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19	27875	28103	28330	28556	28780						23	45	68	90	113	135	158	181	203
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20	30103	30320	30535	30750	30963						21	42	64	85	106	127	148	170	191
21	32222	32428	32634	32838	33041						20	40	61	81	101	121	141	162	182
22	34242	34439	34635	34830	35025						19	39	58	77	97	116	135	155	174
23	36173	36361	36549	36736	36922						18	37	56	74	92	111	130	148	166
24	38021	38202	38382	38561	38739						18	35	53	71	89	106	124	142	160
25	39794	39967	40140	40312	40483						17	34	51	68	85	102	119	136	153
26	41497	41664	41830	41996	42160						16	33	49	66	82	98	115	131	148
27	43136	43297	43457	43616	43775						16	32	47	63	79	95	111	126	142
28	44716	44871	45025	45179	45332						15	30	46	61	76	91	107	122	137
29	46240	46389	46538	46687	46835						15	29	44	59	74	88	103	118	133
30	47712	47857	48001	48144	48287						14	28	43	57	71	85	100	114	128
31	49136	49276	49415	49554	49693						14	28	41	55	69	83	97	110	124
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34	53148	53275	53403	53529	53656						13	25	38	50	63	76	88	101	113
35	54407	54531	54654	54777	54900						12	24	37	49	61	73	86	98	110
36	55630	55751	55871	55991	56110						12	24	36	48	60	71	83	95	107
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39	59106	59218	59329	59439	59550						11	22	33	44	55	66	77	88	99
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42	62325	62428	62531	62634	62737						10	20	31	41	51	61	72	82	92
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62	79239	79309	79379	79449	79518	79588	79657	79727	79796	79865	7	14	21	28	35	42	49	56	63
63	79934	80003	80072	80140	80209	80277	80346	80414	80482	80550	7	14	21	27	34	41	48	55	62
64	80618	80686	80754	80821	80889	80956	81023	81090	81158	81224	7	13	20	27	34	40	47	54	61
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68	83251	83315	83378	83442	83506	83569	83632	83696	83759	83822	6	13	19	25	32	38	44	51	57
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81	90848	90902	90956	91009	91062	91116	91169	91222	91275	91328	5	11	16	21	27	32	37	43	48
82	91381	91434	91487	91540	91593	91645	91698	91751	91803	91855	5	11	16	21	26	32	37	42	47
83	91908	91960	92012	92064	92117	92169	92221	92273	92324	92376	5	10	16	21	26	31	36	42	47
84	92428	92480	92531	92583	92634	92686	92737	92788	92840	92891	5	10	15	21	26	31	36	41	46
85	92942	92993	93044	93095	93146	93197	93247	93298	93349	93399	5	10	15	20	26	31	36	41	46
86	93450	93500	93551	93601	93651	93702	93752	93802	93852	93902	5	10	15	20	25	30	35	40	45
87	93952	94002	94052	94101	94151	94201	94250	94300	94349	94399	5	10	15	20	25	30	35	40	45
88	94448	94498	94547	94596	94645	94694	94743	94792	94841	94890	5	10	15	20	25	29	34	39	44
89	94939	94988	95036	95085	95134	95182	95231	95279	95328	95376	5	10	15	19	24	29	34	39	44
90	95424	95472	95521	95569	95617	95665	95713	95761	95809	95856	5	10	14	19	24	29	34	38	43
91	95904	95952	95999	96047	96095	96142	96190	96237	96284	96332	5	9	14	19	24	28	33	38	43
92	96379	96426	96473	96520	96567	96614	96661	96708	96755	96802	5	9	14	19	24	28	33	38	42
93	96848	96895	96942	96988	97035	97081	97128	97174	97220	97267	5	9	14	19	23	28	33	37	42
94	97313	97359	97405	97451	97497	97543	97589	97635	97681	97727	5	9	14	18	23	28	32	37	42
95	97772	97818	97864	97909	97955	98000	98046	98091	98137	98182	5	9	14	18	23	27	32	36	41
96	98227	98272	98318	98363	98408	98453	98498	98543	98588	98632	5	9	14	18	23	27	32	36	41
97	98677	98722	98767	98811	98856	98900	98945	98989	99034	99078	4	9	13	18	22	27	31	36	40
98	99123	99167	99211	99255	99300	99344	99388	99432	99476	99520	4	9	13	18	22	26	31	35	40
99	99564	99607	99651	99695	99739	99782	99826	99870	99913	99957	4	9	13	17	22	26	31	35	39
	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9

	0	1	2	3	4	5	6	7	8	9	Subtract Differences.									
											1	2	3	4	5	6	7	8	9	
10	1000	9901	9804	9709	9615	9524	9434	9346	9259	9174	Mean differences not sufficiently accurate.									
11	9091	9009	8929	8850	8772	8696	8621	8547	8475	8403										
12	8333	8264	8197	8130	8065	8000	7937	7874	7813	7752										
13	7692	7634	7576	7519	7463	7407	7353	7299	7246	7194										
14	7143	7092	7042	6993	6944	6897	6849	6803	6757	6711										
15	6667	6623	6579	6536	6494	6452	6410	6369	6329	6289	4	8	13	17	21	25	29	33	38	
16	6250	6211	6173	6135	6098	6061	6024	5988	5952	5917	4	7	11	15	18	22	26	29	33	
17	5882	5848	5814	5780	5747	5714	5682	5650	5618	5587	3	6	10	13	16	20	23	26	29	
18	5550	5525	5495	5464	5435	5405	5376	5348	5319	5291	3	6	9	12	15	17	20	23	26	
19	5263	5236	5208	5181	5155	5128	5102	5076	5051	5025	3	5	8	11	13	16	18	21	24	
20	5000	4975	4950	4926	4902	4878	4854	4831	4808	4785	2	5	7	10	12	14	17	19	21	
21	4762	4739	4717	4695	4673	4651	4630	4608	4587	4566	2	4	7	9	11	13	15	17	19	
22	4545	4525	4505	4484	4464	4444	4425	4405	4386	4367	2	4	6	8	10	12	14	16	18	
23	4348	4329	4310	4292	4274	4255	4237	4219	4202	4184	2	4	5	7	9	11	13	14	16	
24	4167	4149	4132	4115	4098	4082	4065	4049	4032	4016	2	3	5	7	8	10	12	13	15	
25	4000	3984	3968	3953	3937	3922	3906	3891	3876	3861	2	3	5	6	8	9	11	12	14	
26	3846	3831	3817	3802	3788	3774	3759	3745	3731	3717	1	3	4	6	7	8	10	11	13	
27	3704	3690	3676	3663	3650	3636	3623	3610	3597	3584	1	3	4	5	7	8	9	11	12	
28	3571	3559	3546	3534	3521	3509	3497	3484	3472	3460	1	2	4	5	6	7	9	10	11	
29	3448	3436	3425	3413	3401	3390	3378	3367	3356	3344	1	2	3	5	6	7	8	9	10	
30	3333	3322	3311	3300	3289	3279	3268	3257	3247	3236	1	2	3	4	5	6	7	9	10	
31	3226	3215	3205	3195	3185	3175	3165	3155	3145	3135	1	2	3	4	5	6	7	8	9	
32	3125	3115	3106	3096	3086	3077	3067	3058	3049	3040	1	2	3	4	5	6	7	8	9	
33	3030	3021	3012	3003	2994	2985	2976	2967	2959	2950	1	2	3	4	4	5	6	7	8	
34	2941	2933	2924	2915	2907	2899	2890	2882	2874	2865	1	2	3	3	4	5	6	7	8	
35	2857	2849	2841	2833	2825	2817	2809	2801	2793	2786	1	2	2	3	4	5	6	6	7	
36	2778	2770	2762	2755	2747	2740	2732	2725	2717	2710	1	2	2	3	4	5	5	6	7	
37	2703	2695	2688	2681	2674	2667	2660	2653	2646	2639	1	1	2	3	4	4	5	6	6	
38	2632	2625	2618	2611	2604	2597	2591	2584	2577	2571	1	1	2	3	3	4	5	5	6	
39	2564	2558	2551	2545	2538	2532	2525	2519	2513	2506	1	1	2	3	3	4	4	5	6	
40	2500	2494	2488	2481	2475	2469	2463	2457	2451	2445	1	1	2	2	3	4	4	5	5	
41	2439	2433	2427	2421	2415	2410	2404	2398	2392	2387	1	1	2	2	3	3	4	5	5	
42	2381	2375	2370	2364	2358	2353	2347	2342	2336	2331	1	1	2	2	3	3	4	4	5	
43	2326	2320	2315	2309	2304	2299	2294	2288	2283	2278	1	1	2	2	3	3	4	4	5	
44	2273	2268	2262	2257	2252	2247	2242	2237	2232	2227	1	1	2	2	3	3	4	4	5	
45	2222	2217	2212	2208	2203	2198	2193	2188	2183	2179	0	1	1	2	2	3	3	4	4	
46	2174	2169	2165	2160	2155	2151	2146	2141	2137	2132	0	1	1	2	2	3	3	4	4	
47	2128	2123	2119	2114	2110	2105	2101	2096	2092	2088	0	1	1	2	2	3	3	4	4	
48	2083	2079	2075	2070	2066	2062	2058	2053	2049	2045	0	1	1	2	2	3	3	3	4	
49	2041	2037	2033	2028	2024	2020	2016	2012	2008	2004	0	1	1	2	2	2	3	3	4	
50	2000	1996	1992	1988	1984	1980	1976	1972	1969	1965	0	1	1	2	2	2	3	3	4	
51	1961	1957	1953	1949	1946	1942	1938	1934	1931	1927	0	1	1	2	2	2	3	3	3	
52	1923	1919	1916	1912	1908	1905	1901	1898	1894	1890	0	1	1	1	2	2	3	3	3	
53	1887	1883	1880	1876	1873	1869	1866	1862	1859	1855	0	1	1	1	2	2	2	3	3	
54	1852	1848	1845	1842	1838	1835	1832	1828	1825	1821	0	1	1	1	2	2	2	3	3	
	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9	
												Subtract Differences.								

	0	1	2	3	4	5	6	7	8	9	Subtract Differences.								
											1	2	3	4	5	6	7	8	9
55	1818	1815	1812	1808	1805	1802	1799	1795	1792	1789	0	1	1	1	2	2	2	3	3
56	1786	1783	1779	1776	1773	1770	1767	1764	1761	1757	0	1	1	1	2	2	2	3	3
57	1754	1751	1748	1745	1742	1739	1736	1733	1730	1727	0	1	1	1	2	2	2	2	3
58	1724	1721	1718	1715	1712	1709	1706	1704	1701	1698	0	1	1	1	1	2	2	2	3
59	1695	1692	1689	1686	1684	1681	1678	1675	1672	1669	0	1	1	1	1	2	2	2	3
60	1667	1664	1661	1658	1656	1653	1650	1647	1645	1642	0	1	1	1	1	2	2	2	3
61	1639	1637	1634	1631	1629	1626	1623	1621	1618	1616	0	1	1	1	1	2	2	2	2
62	1613	1610	1608	1605	1603	1600	1597	1595	1592	1590	0	1	1	1	1	2	2	2	2
63	1587	1585	1582	1580	1577	1575	1572	1570	1567	1565	0	0	1	1	1	1	2	2	2
64	1563	1560	1558	1555	1553	1550	1548	1546	1543	1541	0	0	1	1	1	1	2	2	2
65	1538	1536	1534	1531	1529	1527	1524	1522	1520	1517	0	0	1	1	1	1	2	2	2
66	1515	1513	1511	1508	1506	1504	1502	1499	1497	1495	0	0	1	1	1	1	2	2	2
67	1493	1490	1488	1486	1484	1481	1479	1477	1475	1473	0	0	1	1	1	1	2	2	2
68	1471	1468	1466	1464	1462	1460	1458	1456	1453	1451	0	0	1	1	1	1	2	2	2
69	1449	1447	1445	1443	1441	1439	1437	1435	1433	1431	0	0	1	1	1	1	1	2	2
70	1429	1427	1425	1422	1420	1418	1416	1414	1412	1410	0	0	1	1	1	1	1	2	2
71	1408	1406	1404	1403	1401	1399	1397	1395	1393	1391	0	0	1	1	1	1	1	2	2
72	1389	1387	1385	1383	1381	1379	1377	1376	1374	1372	0	0	1	1	1	1	1	2	2
73	1370	1368	1366	1364	1362	1361	1359	1357	1355	1353	0	0	1	1	1	1	1	2	2
74	1351	1350	1348	1346	1344	1342	1340	1339	1337	1335	0	0	1	1	1	1	1	1	2
75	1333	1332	1330	1328	1326	1325	1323	1321	1319	1318	0	0	1	1	1	1	1	1	2
76	1316	1314	1312	1311	1309	1307	1305	1304	1302	1300	0	0	1	1	1	1	1	1	2
77	1299	1297	1295	1294	1292	1290	1289	1287	1285	1284	0	0	0	1	1	1	1	1	1
78	1282	1280	1279	1277	1276	1274	1272	1271	1269	1267	0	0	0	1	1	1	1	1	1
79	1266	1264	1263	1261	1259	1258	1256	1255	1253	1252	0	0	0	1	1	1	1	1	1
80	1250	1248	1247	1245	1244	1242	1241	1239	1238	1236	0	0	0	1	1	1	1	1	1
81	1235	1233	1232	1230	1229	1227	1225	1224	1222	1221	0	0	0	1	1	1	1	1	1
82	1220	1218	1217	1215	1214	1212	1211	1209	1208	1206	0	0	0	1	1	1	1	1	1
83	1205	1203	1202	1200	1199	1198	1196	1195	1193	1192	0	0	0	1	1	1	1	1	1
84	1190	1189	1188	1186	1185	1183	1182	1181	1179	1178	0	0	0	1	1	1	1	1	1
85	1176	1175	1174	1172	1171	1170	1168	1167	1166	1164	0	0	0	1	1	1	1	1	1
86	1163	1161	1160	1159	1157	1156	1155	1153	1152	1151	0	0	0	1	1	1	1	1	1
87	1149	1148	1147	1145	1144	1143	1142	1140	1139	1138	0	0	0	1	1	1	1	1	1
88	1136	1135	1134	1133	1131	1130	1129	1127	1126	1125	0	0	0	1	1	1	1	1	1
89	1124	1122	1121	1120	1119	1117	1116	1115	1114	1112	0	0	0	1	1	1	1	1	1
90	1111	1110	1109	1107	1106	1105	1104	1103	1101	1100	0	0	0	1	1	1	1	1	1
91	1099	1098	1096	1095	1094	1093	1092	1091	1089	1088	0	0	0	0	1	1	1	1	1
92	1087	1086	1085	1083	1082	1081	1080	1079	1078	1076	0	0	0	0	1	1	1	1	1
93	1075	1074	1073	1072	1071	1070	1068	1067	1066	1065	0	0	0	0	1	1	1	1	1
94	1064	1063	1062	1060	1059	1058	1057	1056	1055	1054	0	0	0	0	1	1	1	1	1
95	1053	1052	1050	1049	1048	1047	1046	1045	1044	1043	0	0	0	0	1	1	1	1	1
96	1042	1041	1040	1038	1037	1036	1035	1034	1033	1032	0	0	0	0	1	1	1	1	1
97	1031	1030	1029	1028	1027	1026	1025	1024	1022	1021	0	0	0	0	1	1	1	1	1
98	1020	1019	1018	1017	1016	1015	1014	1013	1012	1011	0	0	0	0	1	1	1	1	1
99	1010	1009	1008	1007	1006	1005	1004	1003	1002	1001	0	0	0	0	0	1	1	1	1
	0	1	2	3	4	5	6	7	8	9	Subtract Differences.								
											1	2	3	4	5	6	7	8	9

SQUARES

	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9
1·0	1'000	1'020	1'040	1'061	1'082	1'103	1'124	1'145	1'166	1'188	2	4	6	8	10	13	15	17	19
1·1	1'210	1'232	1'254	1'277	1'300	1'323	1'346	1'369	1'392	1'416	2	5	7	9	11	14	16	18	21
1·2	1'440	1'464	1'488	1'513	1'538	1'563	1'588	1'613	1'638	1'664	2	5	7	10	12	15	17	20	22
1·3	1'690	1'716	1'742	1'769	1'796	1'823	1'850	1'877	1'904	1'932	3	5	8	11	13	16	19	22	24
1·4	1'960	1'988	2'016	2'045	2'074	2'103	2'132	2'161	2'190	2'220	3	6	9	12	14	17	20	23	26
1·5	2'250	2'280	2'310	2'341	2'372	2'403	2'434	2'465	2'496	2'528	3	6	9	12	15	19	22	25	28
1·6	2'560	2'592	2'624	2'657	2'690	2'723	2'756	2'789	2'822	2'856	3	7	10	13	16	20	23	26	30
1·7	2'800	2'924	2'958	2'993	3'028	3'063	3'098	3'133	3'168	3'204	3	7	10	14	17	21	24	28	31
1·8	3'240	3'276	3'312	3'349	3'386	3'423	3'460	3'497	3'534	3'572	4	7	11	15	18	22	26	30	33
1·9	3'610	3'648	3'686	3'725	3'764	3'803	3'842	3'881	3'920	3'960	4	8	12	16	19	23	27	31	35
2·0	4'000	4'040	4'080	4'121	4'162	4'203	4'244	4'285	4'326	4'368	4	8	12	16	20	25	29	33	37
2·1	4'410	4'452	4'494	4'537	4'580	4'623	4'666	4'709	4'752	4'796	4	9	13	17	21	26	30	34	39
2·2	4'840	4'884	4'928	4'973	5'018	5'063	5'108	5'153	5'198	5'244	4	9	13	18	22	27	31	36	40
2·3	5'290	5'336	5'382	5'429	5'476	5'523	5'570	5'617	5'664	5'712	5	9	14	19	23	28	33	38	42
2·4	5'760	5'808	5'856	5'905	5'954	6'003	6'052	6'101	6'150	6'200	5	10	15	20	24	29	34	39	44
2·5	6'250	6'300	6'350	6'401	6'452	6'503	6'554	6'605	6'656	6'708	5	10	15	20	25	31	36	41	46
2·6	6'760	6'812	6'864	6'917	6'970	7'023	7'076	7'129	7'182	7'236	5	11	16	21	26	32	37	42	48
2·7	7'290	7'344	7'398	7'453	7'508	7'563	7'618	7'673	7'728	7'784	5	11	16	22	27	33	38	44	49
2·8	7'840	7'896	7'952	8'009	8'066	8'123	8'180	8'237	8'294	8'352	6	11	17	23	28	34	40	46	51
2·9	8'410	8'468	8'526	8'585	8'644	8'703	8'762	8'821	8'880	8'940	6	12	18	24	29	35	41	47	53
3·0	9'000	9'060	9'120	9'181	9'242	9'303	9'364	9'425	9'486	9'548	6	12	18	24	30	37	43	49	55
3·1	9'610	9'672	9'734	9'797	9'860	9'923	9'986	10'05	10'11	10'18	6	13	19	25	31	38	44	50	57
3·2	10'24	10'30	10'37	10'43	10'50	10'56	10'63	10'69	10'76	10'82	1	1	2	3	3	4	5	5	6
3·3	10'89	10'96	11'02	11'09	11'16	11'22	11'29	11'36	11'42	11'49	1	1	2	3	3	4	5	5	6
3·4	11'56	11'63	11'70	11'76	11'83	11'90	11'97	12'04	12'11	12'18	1	1	2	3	3	4	5	5	6
3·5	12'25	12'32	12'39	12'46	12'53	12'60	12'67	12'74	12'82	12'89	1	1	2	3	4	4	5	6	6
3·6	12'96	13'03	13'10	13'18	13'25	13'32	13'40	13'47	13'54	13'62	1	1	2	3	4	4	5	6	7
3·7	13'69	13'76	13'84	13'91	13'99	14'06	14'14	14'21	14'29	14'36	1	2	2	3	4	4	5	6	7
3·8	14'44	14'52	14'59	14'67	14'75	14'82	14'90	14'98	15'05	15'13	1	2	2	3	4	4	5	6	7
3·9	15'21	15'29	15'37	15'44	15'52	15'60	15'68	15'76	15'84	15'92	1	2	2	3	4	4	5	6	7
4·0	16'00	16'08	16'16	16'24	16'32	16'40	16'48	16'56	16'65	16'73	1	2	2	3	4	4	5	6	7
4·1	16'81	16'89	16'97	17'06	17'14	17'22	17'31	17'39	17'47	17'56	1	2	2	3	4	4	5	6	7
4·2	17'64	17'72	17'81	17'89	17'98	18'06	18'15	18'23	18'32	18'40	1	2	3	3	4	4	5	6	7
4·3	18'49	18'58	18'66	18'75	18'84	18'92	19'01	19'10	19'18	19'27	1	2	3	3	4	4	5	6	7
4·4	19'36	19'45	19'54	19'62	19'71	19'80	19'89	19'98	20'07	20'16	1	2	3	4	4	4	5	6	7
4·5	20'25	20'34	20'43	20'52	20'61	20'70	20'79	20'88	20'98	21'07	1	2	3	4	5	5	6	7	8
4·6	21'16	21'25	21'34	21'44	21'53	21'62	21'72	21'81	21'90	22'00	1	2	3	4	5	5	6	7	8
4·7	22'09	22'18	22'28	22'37	22'47	22'56	22'66	22'75	22'85	22'94	1	2	3	4	5	5	6	7	8
4·8	23'04	23'14	23'23	23'33	23'43	23'52	23'62	23'72	23'81	23'91	1	2	3	4	5	5	6	7	8
4·9	24'01	24'11	24'21	24'30	24'40	24'50	24'60	24'70	24'80	24'90	1	2	3	4	5	5	6	7	8
5·0	25'00	25'10	25'20	25'30	25'40	25'50	25'60	25'70	25'81	25'91	1	2	3	4	5	5	6	7	8
5·1	26'01	26'11	26'21	26'32	26'42	26'52	26'63	26'73	26'83	26'94	1	2	3	4	5	5	6	7	8
5·2	27'04	27'14	27'25	27'35	27'46	27'56	27'67	27'77	27'88	27'98	1	2	3	4	5	5	6	7	8
5·3	28'09	28'20	28'30	28'41	28'52	28'62	28'73	28'84	28'94	29'05	1	2	3	4	5	5	6	7	9
5·4	29'16	29'27	29'38	29'48	29'59	29'70	29'81	29'92	30'03	30'14	1	2	3	4	5	5	6	7	10
	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9

	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9
5·5	30'25	30'36	30'47	30'58	30'69	30'80	30'91	31'02	31'14	31'25	1	2	3	4	5	6	7	8	9 10
5·6	31'36	31'47	31'58	31'70	31'81	31'92	32'04	32'15	32'26	32'38	1	2	3	5	6	7	8	9 10	
5·7	32'49	32'60	32'72	32'83	32'95	33'06	33'18	33'29	33'41	33'52	1	2	3	5	6	7	8	9 10	
5·8	33'64	33'76	33'87	33'99	34'11	34'22	34'34	34'46	34'57	34'69	1	2	4	5	6	7	8	9 11	
5·9	34'81	34'93	35'05	35'16	35'28	35'40	35'52	35'64	35'76	35'88	1	2	4	5	6	7	8	10 11	
6·0	36'00	36'12	36'24	36'36	36'48	36'60	36'72	36'84	36'97	37'09	1	2	4	5	6	7	8	10 11	
6·1	37'21	37'33	37'45	37'58	37'70	37'82	37'95	38'07	38'19	38'32	1	2	4	5	6	7	9	10 11	
6·2	38'44	38'56	38'69	38'81	38'94	39'06	39'19	39'31	39'44	39'56	1	3	4	5	6	8	9	10 11	
6·3	39'69	39'82	39'94	40'07	40'20	40'32	40'45	40'58	40'70	40'83	1	3	4	5	6	8	9	10 11	
6·4	40'96	41'09	41'22	41'34	41'47	41'60	41'73	41'86	41'99	42'12	1	3	4	5	6	8	9	10 12	
6·5	42'25	42'38	42'51	42'64	42'77	42'90	43'03	43'16	43'30	43'43	1	3	4	5	7	8	9	10 12	
6·6	43'56	43'69	43'82	43'96	44'09	44'22	44'36	44'49	44'62	44'76	1	3	4	5	7	8	9	11 12	
6·7	44'89	45'02	45'16	45'29	45'43	45'56	45'70	45'83	45'97	46'10	1	3	4	5	7	8	9	11 12	
6·8	46'24	46'38	46'51	46'65	46'79	46'92	47'06	47'20	47'33	47'47	1	3	4	5	7	8	10 11 12		
6·9	47'61	47'75	47'89	48'02	48'16	48'30	48'44	48'58	48'72	48'86	1	3	4	6	7	8	10 11 13		
7·0	49'00	49'14	49'28	49'42	49'56	49'70	49'84	49'98	50'13	50'27	1	3	4	6	7	8	10 11 13		
7·1	50'41	50'55	50'69	50'84	50'98	51'12	51'27	51'41	51'55	51'70	1	3	4	6	7	9	10 11 13		
7·2	51'84	51'98	52'13	52'27	52'42	52'56	52'71	52'85	53'00	53'14	1	3	4	6	7	9	10 12 13		
7·3	53'29	53'44	53'58	53'73	53'88	54'02	54'17	54'32	54'46	54'61	1	3	4	6	7	9	10 12 13		
7·4	54'76	54'91	55'06	55'20	55'35	55'50	55'65	55'80	55'95	56'10	1	3	4	6	7	9	10 12 13		
7·5	56'25	56'40	56'55	56'70	56'85	57'00	57'15	57'30	57'46	57'61	2	3	5	6	8	9	11 12 14		
7·6	57'76	57'91	58'06	58'22	58'37	58'52	58'68	58'83	58'98	59'14	2	3	5	6	8	9	11 12 14		
7·7	59'29	59'44	59'60	59'75	59'91	60'06	60'22	60'37	60'53	60'68	2	3	5	6	8	9	11 12 14		
7·8	60'84	61'00	61'15	61'31	61'47	61'62	61'78	61'94	62'09	62'25	2	3	5	6	8	9	11 13 14		
7·9	62'41	62'57	62'73	62'88	63'04	63'20	63'36	63'52	63'68	63'84	2	3	5	6	8	10 11 13 14			
8·0	64'00	64'16	64'32	64'48	64'64	64'80	64'96	65'12	65'29	65'45	2	3	5	6	8	10 11 13 14			
8·1	65'61	65'77	65'93	66'10	66'26	66'42	66'59	66'75	66'91	67'08	2	3	5	7	8	10 11 13 15			
8·2	67'24	67'40	67'57	67'73	67'90	68'06	68'23	68'39	68'56	68'72	2	3	5	7	8	10 12 13 15			
8·3	68'89	69'06	69'22	69'39	69'56	69'72	69'89	70'06	70'22	72'39	2	3	5	7	8	10 12 13 15			
8·4	70'56	70'73	70'90	71'06	71'23	71'40	71'57	71'74	71'91	72'08	2	3	5	7	8	10 12 14 15			
8·5	72'25	72'42	72'59	72'76	72'93	73'10	73'27	73'44	73'62	73'79	2	3	5	7	9	10 12 14 15			
8·6	73'96	74'13	74'30	74'48	74'65	75'82	75'00	75'17	75'34	75'52	2	3	5	7	9	10 12 14 16			
8·7	75'69	75'86	76'04	76'21	76'39	76'56	76'74	76'91	77'09	77'26	2	4	5	7	9	11 12 14 16			
8·8	77'44	77'62	77'79	77'97	78'15	78'32	78'50	78'68	78'85	79'03	2	4	5	7	9	11 12 14 16			
8·9	79'21	79'39	79'57	79'74	79'92	80'10	80'28	80'46	80'64	80'82	2	4	5	7	9	11 13 14 16			
9·0	81'00	81'18	81'36	81'54	81'72	81'90	82'08	82'26	82'45	82'63	2	4	5	7	9	11 13 14 16			
9·1	82'81	82'99	83'17	83'36	83'54	83'72	83'91	84'09	84'27	84'46	2	4	5	7	9	11 13 15 16			
9·2	84'64	84'82	85'01	85'19	85'38	85'56	85'75	85'93	86'12	86'30	2	4	6	7	9	11 13 15 17			
9·3	86'49	86'68	86'86	87'05	87'24	87'42	87'61	87'80	87'98	88'17	2	4	6	7	9	11 13 15 17			
9·4	88'36	88'55	88'74	88'92	89'11	89'30	89'49	89'68	89'87	90'06	2	4	6	8	9	11 13 15 17			
9·5	90'25	90'44	90'63	90'82	91'01	91'20	91'39	91'58	91'78	91'97	2	4	6	8	10	11 13 15 17			
9·6	92'16	92'35	92'54	92'74	92'93	93'12	93'32	93'51	93'70	93'90	2	4	6	8	10	12 14 15 17			
9·7	94'09	94'28	94'48	94'67	94'87	95'06	95'26	95'45	95'65	95'84	2	4	6	8	10	12 14 16 18			
9·8	96'04	96'24	96'43	96'63	96'83	97'02	97'22	97'42	97'61	97'81	2	4	6	8	10	12 14 16 18			
9·9	98'01	98'21	98'41	98'60	98'80	99'00	99'20	99'40	99'60	99'80	2	4	6	8	10	12 14 16 18			
	0	1	2	3	4	5	6	7	8	9	1	2	3	4	5	6	7	8	9

NATURAL SINES

	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'
0°	'0000	'0017	'0035	'0052	'0070	'0087	'0105	'0122	'0140	'0157	3	6	9	12	15
1	'0175	0192	0209	0227	0244	0262	0279	0297	0314	0332	3	6	9	12	15
2	'0349	0366	0384	0401	0419	0436	0454	0471	0488	0506	3	6	9	12	15
3	'0523	0541	0558	0576	0593	0610	0628	0645	0663	0680	3	6	9	12	15
4	'0698	0715	0732	0750	0767	0785	0802	0819	0837	0854	3	6	9	12	14
5	'0872	0889	0906	0924	0941	0958	0976	0993	1011	1028	3	6	9	12	14
6	'1045	1063	1080	1097	1115	1132	1149	1167	1184	1201	3	6	9	12	14
7	'1219	1236	1253	1271	1288	1305	1323	1340	1357	1374	3	6	9	12	14
8	'1392	1409	1426	1444	1461	1478	1495	1513	1530	1547	3	6	9	12	14
9	'1564	1582	1599	1616	1633	1650	1668	1685	1702	1719	3	6	9	11	14
10	'1736	1754	1771	1788	1805	1822	1840	1857	1874	1891	3	6	9	11	14
11	'1908	1925	1942	1959	1977	1994	2011	2028	2045	2062	3	6	9	11	14
12	'2079	2096	2113	2130	2147	2164	2181	2198	2215	2233	3	6	9	11	14
13	'2250	2267	2284	2300	2317	2334	2351	2368	2385	2402	3	6	8	11	14
14	'2419	2436	2453	2470	2487	2504	2521	2538	2554	2571	3	6	8	11	14
15	'2588	2605	2622	2639	2656	2672	2689	2706	2723	2740	3	6	8	11	14
16	'2756	2773	2790	2807	2823	2840	2857	2874	2890	2907	3	6	8	11	14
17	'2924	2940	2957	2974	2990	3007	3024	3040	3057	3074	3	6	8	11	14
18	'3090	3107	3123	3140	3156	3173	3190	3206	3223	3239	3	6	8	11	14
19	'3256	3272	3289	3305	3322	3338	3355	3371	3387	3404	3	5	8	11	14
20	'3420	3437	3453	3469	3486	3502	3518	3535	3551	3567	3	5	8	11	14
21	'3584	3600	3616	3633	3649	3665	3681	3697	3714	3730	3	5	8	11	14
22	'3746	3762	3778	3795	3811	3827	3843	3859	3875	3891	3	5	8	11	13
23	'3907	3923	3939	3955	3971	3987	4003	4019	4035	4051	3	5	8	11	13
24	'4067	4083	4099	4115	4131	4147	4163	4179	4195	4210	3	5	8	11	13
25	'4226	4242	4258	4274	4289	4305	4321	4337	4352	4368	3	5	8	11	13
26	'4384	4399	4415	4431	4446	4462	4478	4493	4509	4524	3	5	8	10	13
27	'4540	4555	4571	4586	4602	4617	4633	4648	4664	4679	3	5	8	10	13
28	'4695	4710	4726	4741	4756	4772	4787	4802	4818	4833	3	5	8	10	13
29	'4848	4863	4879	4894	4909	4924	4939	4955	4970	4985	3	5	8	10	13
30	'5000	5015	5030	5045	5060	5075	5090	5105	5120	5135	3	5	8	10	13
31	'5150	5165	5180	5195	5210	5225	5240	5255	5270	5284	2	5	7	10	12
32	'5299	5314	5329	5344	5358	5373	5388	5402	5417	5432	2	5	7	10	12
33	'5446	5461	5476	5490	5505	5519	5534	5548	5563	5577	2	5	7	10	12
34	'5592	5606	5621	5635	5650	5664	5678	5693	5707	5721	2	5	7	10	12
35	'5736	5750	5764	5779	5793	5807	5821	5835	5850	5864	2	5	7	9	12
36	'5878	5892	5906	5920	5934	5948	5962	5976	5990	6004	2	5	7	9	12
37	'6018	6032	6046	6060	6074	6088	6101	6115	6129	6143	2	5	7	9	12
38	'6157	6170	6184	6198	6211	6225	6239	6252	6266	6280	2	5	7	9	11
39	'6293	6307	6320	6334	6347	6361	6374	6388	6401	6414	2	4	7	9	11
40	'6428	6441	6455	6468	6481	6494	6508	6521	6534	6547	2	4	7	9	11
41	'6561	6574	6587	6600	6613	6626	6639	6652	6665	6678	2	4	7	9	11
42	'6691	6704	6717	6730	6743	6756	6769	6782	6794	6807	2	4	6	9	11
43	'6820	6833	6845	6858	6871	6884	6896	6909	6921	6934	2	4	6	8	11
44	'6947	6959	6972	6984	6997	7009	7022	7034	7046	7059	2	4	6	8	10
	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'

	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'
45°	7071	7083	7096	7108	7120	7133	7145	7157	7169	7181	2	4	6	8	10
46	7193	7206	7218	7230	7242	7254	7266	7278	7290	7302	2	4	6	8	10
47	7314	7325	7337	7349	7361	7373	7385	7396	7408	7420	2	4	6	8	10
48	7431	7443	7455	7466	7478	7490	7501	7513	7524	7536	2	4	6	8	10
49	7547	7559	7570	7581	7593	7604	7615	7627	7638	7649	2	4	6	8	9
50	7660	7672	7683	7694	7705	7716	7727	7738	7749	7760	2	4	6	7	9
51	7771	7782	7793	7804	7815	7826	7837	7848	7859	7869	2	4	5	7	9
52	7880	7891	7902	7912	7923	7934	7944	7955	7965	7976	2	4	5	7	9
53	7986	7997	8007	8018	8028	8039	8049	8059	8070	8080	2	3	5	7	9
54	8090	8100	8111	8121	8131	8141	8151	8161	8171	8181	2	3	5	7	8
55	8192	8202	8211	8221	8231	8241	8251	8261	8271	8281	2	3	5	7	8
56	8290	8300	8310	8320	8329	8339	8348	8358	8368	8377	2	3	5	6	8
57	8387	8396	8406	8415	8425	8434	8443	8453	8462	8471	2	3	5	6	8
58	8480	8490	8499	8508	8517	8526	8536	8545	8554	8563	2	3	5	6	8
59	8572	8581	8590	8599	8607	8616	8625	8634	8643	8652	1	3	4	6	7
60	8660	8669	8678	8686	8695	8704	8712	8721	8729	8738	1	3	4	6	7
61	8746	8755	8763	8771	8780	8788	8796	8805	8813	8821	1	3	4	6	7
62	8829	8838	8846	8854	8862	8870	8878	8886	8894	8902	1	3	4	5	7
63	8910	8918	8926	8934	8942	8949	8957	8965	8973	8980	1	3	4	5	6
64	8988	8996	9003	9011	9018	9026	9033	9041	9048	9056	1	3	4	5	6
65	9063	9070	9078	9085	9092	9100	9107	9114	9121	9128	1	2	4	5	6
66	9135	9143	9150	9157	9164	9171	9178	9184	9191	9198	1	2	3	5	6
67	9205	9212	9219	9225	9232	9239	9245	9252	9259	9265	1	2	3	4	6
68	9272	9278	9285	9291	9298	9304	9311	9317	9323	9330	1	2	3	4	5
69	9336	9342	9348	9354	9361	9367	9373	9379	9385	9391	1	2	3	4	5
70	9397	9403	9409	9415	9421	9426	9432	9438	9444	9449	1	2	3	4	5
71	9455	9461	9466	9472	9478	9483	9489	9494	9500	9505	1	2	3	4	5
72	9511	9516	9521	9527	9532	9537	9542	9548	9553	9558	1	2	3	3	4
73	9563	9568	9573	9578	9583	9588	9593	9598	9603	9608	1	2	2	3	4
74	9613	9617	9622	9627	9632	9636	9641	9646	9650	9655	1	2	2	3	4
75	9659	9664	9668	9673	9677	9681	9686	9690	9694	9699	1	1	2	3	4
76	9703	9707	9711	9715	9720	9724	9728	9732	9736	9740	1	1	2	3	3
77	9744	9748	9751	9755	9759	9763	9767	9770	9774	9778	1	1	2	3	3
78	9781	9785	9789	9792	9796	9799	9803	9806	9810	9813	1	1	2	2	3
79	9816	9820	9823	9826	9829	9833	9836	9839	9842	9845	1	1	2	2	3
80	9848	9851	9854	9857	9860	9863	9866	9869	9871	9874	0	1	1	2	2
81	9877	9880	9882	9885	9888	9890	9893	9895	9898	9900	0	1	1	2	2
82	9903	9905	9907	9910	9912	9914	9917	9919	9921	9923	0	1	1	2	2
83	9925	9928	9930	9932	9934	9936	9938	9940	9942	9943	0	1	1	1	2
84	9945	9947	9949	9951	9952	9954	9956	9957	9959	9960	0	1	1	1	1
85	9962	9963	9965	9966	9968	9969	9971	9972	9973	9974	0	0	1	1	1
86	9976	9977	9978	9979	9980	9981	9982	9983	9984	9985	0	0	1	1	1
87	9986	9987	9988	9989	9990	9990	9991	9992	9993	9993	0	0	0	1	1
88	9994	9995	9995	9996	9996	9997	9997	9997	9998	9998	0	0	0	0	0
89	9998	9999	9999	9999	9999	1'000	1'000	1'000	1'000	1'000	0	0	0	0	0
	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'

NATURAL COSINES

	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	Subtract Differences.				
											1'	2'	3'	4'	5'
0°	1'000	1'000	1'000	1'000	1'000	1'000	9999	9999	9999	9999	0	0	0	0	0
1	9998	9998	9998	9997	9997	9997	9996	9996	9995	9995	0	0	0	0	0
2	9994	9993	9993	9992	9991	9990	9990	9989	9988	9987	0	0	0	1	1
3	9986	9985	9984	9983	9982	9981	9980	9979	9978	9977	0	0	1	1	1
4	9976	9974	9973	9972	9971	9969	9968	9966	9965	9963	0	0	1	1	1
5	9962	9960	9959	9957	9956	9954	9952	9951	9949	9947	0	1	1	1	1
6	9945	9943	9942	9940	9938	9936	9934	9932	9930	9928	0	1	1	1	2
7	9925	9923	9921	9919	9917	9914	9912	9910	9907	9905	0	1	1	2	2
8	9903	9900	9898	9895	9893	9890	9888	9885	9882	9880	0	1	1	2	2
9	9877	9874	9871	9869	9866	9863	9860	9857	9854	9851	0	1	1	2	2
10	9848	9845	9842	9839	9836	9833	9829	9826	9823	9820	1	1	2	2	3
11	9816	9813	9810	9806	9803	9799	9796	9792	9789	9785	1	1	2	2	3
12	9781	9778	9774	9770	9767	9763	9759	9755	9751	9748	1	1	2	3	3
13	9744	9740	9736	9732	9728	9724	9720	9715	9711	9707	1	1	2	3	3
14	9703	9699	9694	9690	9686	9681	9677	9673	9668	9664	1	1	2	3	4
15	9659	9655	9650	9646	9641	9636	9632	9627	9622	9617	1	2	2	3	4
16	9613	9608	9603	9598	9593	9588	9583	9578	9573	9568	1	2	2	3	4
17	9563	9558	9553	9548	9542	9537	9532	9527	9521	9516	1	2	3	3	4
18	9511	9505	9500	9494	9489	9483	9478	9472	9466	9461	1	2	3	4	5
19	9455	9449	9444	9438	9432	9426	9421	9415	9409	9403	1	2	3	4	5
20	9397	9391	9385	9379	9373	9367	9361	9354	9348	9342	1	2	3	4	5
21	9336	9330	9323	9317	9311	9304	9298	9291	9285	9278	1	2	3	4	5
22	9272	9265	9259	9252	9245	9239	9232	9225	9219	9212	1	2	3	4	6
23	9205	9198	9191	9184	9178	9171	9164	9157	9150	9143	1	2	3	5	6
24	9135	9128	9121	9114	9107	9100	9092	9085	9078	9070	1	2	4	5	6
25	9063	9056	9048	9041	9033	9026	9018	9011	9003	8996	1	3	4	5	6
26	8988	8980	8973	8965	8957	8949	8942	8934	8926	8918	1	3	4	5	6
27	8910	8902	8894	8886	8878	8870	8862	8854	8846	8838	1	3	4	5	7
28	8820	8811	8813	8805	8796	8788	8780	8771	8763	8755	1	3	4	6	7
29	8746	8738	8729	8721	8712	8704	8695	8686	8678	8669	1	3	4	6	7
30	8660	8652	8643	8634	8625	8616	8607	8599	8590	8581	1	3	4	6	7
31	8572	8563	8554	8545	8536	8526	8517	8508	8499	8490	2	3	5	6	8
32	8480	8471	8462	8453	8443	8434	8425	8415	8406	8396	2	3	5	6	8
33	8387	8377	8368	8358	8348	8339	8329	8320	8310	8300	2	3	5	6	8
34	8290	8281	8271	8261	8251	8241	8231	8221	8211	8202	2	3	5	7	8
35	8192	8181	8171	8161	8151	8141	8131	8121	8111	8100	2	3	5	7	8
36	8090	8080	8070	8059	8049	8039	8028	8018	8007	7997	2	3	5	7	9
37	7986	7976	7965	7955	7944	7934	7923	7912	7902	7891	2	4	5	7	9
38	7880	7869	7859	7848	7837	7826	7815	7804	7793	7782	2	4	5	7	9
39	7771	7760	7749	7738	7727	7716	7705	7694	7683	7672	2	4	6	7	9
40	7660	7649	7638	7627	7615	7604	7593	7581	7570	7559	2	4	6	8	9
41	7547	7536	7524	7513	7501	7490	7478	7466	7455	7443	2	4	6	8	10
42	7431	7420	7408	7396	7385	7373	7361	7349	7337	7325	2	4	6	8	10
43	7314	7302	7290	7278	7266	7254	7242	7230	7218	7206	2	4	6	8	10
44	7193	7181	7169	7157	7145	7133	7120	7108	7096	7083	2	4	6	8	10
	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	Subtract Differences.				
											1'	2'	3'	4'	5'

NATURAL COSINES

	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	Subtract Differences.				
											1'	2'	3'	4'	5'
45°	·7071	·7059	·7046	·7034	·7022	·7009	·6997	·6984	·6972	·6959	2	4	6	8	10
46	·6947	·6934	·6921	·6909	·6896	·6884	·6871	·6858	·6845	·6833	2	4	6	8	11
47	·6820	·6807	·6794	·6782	·6769	·6756	·6743	·6730	·6717	·6704	2	4	6	9	11
48	·6691	·6678	·6665	·6652	·6639	·6626	·6613	·6600	·6587	·6574	2	4	7	9	11
49	·6561	·6547	·6534	·6521	·6508	·6494	·6481	·6468	·6455	·6441	2	4	7	9	11
50	·6428	·6414	·6401	·6388	·6374	·6361	·6347	·6334	·6320	·6307	2	4	7	9	11
51	·6293	·6280	·6266	·6252	·6239	·6225	·6211	·6198	·6184	·6170	2	5	7	9	11
52	·6157	·6143	·6129	·6115	·6101	·6088	·6074	·6060	·6046	·6032	2	5	7	9	12
53	·6018	·6004	·5990	·5976	·5962	·5948	·5934	·5920	·5906	·5892	2	5	7	9	12
54	·5878	·5864	·5850	·5835	·5821	·5807	·5793	·5779	·5764	·5750	2	5	7	9	12
55	·5736	·5721	·5707	·5693	·5678	·5664	·5650	·5635	·5621	·5606	2	5	7	10	12
56	·5592	·5577	·5563	·5548	·5534	·5519	·5505	·5490	·5476	·5461	2	5	7	10	12
57	·5446	·5432	·5417	·5402	·5388	·5373	·5358	·5344	·5329	·5314	2	5	7	10	12
58	·5299	·5284	·5270	·5255	·5240	·5225	·5210	·5195	·5180	·5165	2	5	7	10	12
59	·5150	·5135	·5120	·5105	·5090	·5075	·5060	·5045	·5030	·5015	3	5	8	10	13
60	·5000	·4985	·4970	·4955	·4939	·4924	·4909	·4894	·4879	·4863	3	5	8	10	13
61	·4848	·4833	·4818	·4802	·4787	·4772	·4756	·4741	·4726	·4710	3	5	8	10	13
62	·4695	·4679	·4664	·4648	·4633	·4617	·4602	·4586	·4571	·4555	3	5	8	10	13
63	·4540	·4524	·4509	·4493	·4478	·4462	·4446	·4431	·4415	·4399	3	5	8	10	13
64	·4384	·4368	·4352	·4337	·4321	·4305	·4289	·4274	·4258	·4242	3	5	8	11	13
65	·4226	·4210	·4195	·4179	·4163	·4147	·4131	·4115	·4099	·4083	3	5	8	11	13
66	·4067	·4051	·4035	·4019	·4003	·3987	·3971	·3955	·3939	·3923	3	5	8	11	14
67	·3907	·3891	·3875	·3859	·3843	·3827	·3811	·3795	·3778	·3762	3	5	8	11	14
68	·3746	·3730	·3714	·3697	·3681	·3665	·3649	·3633	·3616	·3600	3	5	8	11	14
69	·3584	·3567	·3551	·3535	·3518	·3502	·3486	·3469	·3453	·3437	3	5	8	11	14
70	·3420	·3404	·3387	·3371	·3355	·3338	·3322	·3305	·3289	·3272	3	5	8	11	14
71	·3256	·3239	·3223	·3206	·3190	·3173	·3156	·3140	·3123	·3107	3	6	8	11	14
72	·3090	·3074	·3057	·3040	·3024	·3007	·2990	·2974	·2957	·2940	3	6	8	11	14
73	·2924	·2907	·2890	·2874	·2857	·2840	·2823	·2807	·2790	·2773	3	6	8	11	14
74	·2756	·2740	·2723	·2706	·2689	·2672	·2656	·2639	·2622	·2605	3	6	8	11	14
75	·2588	·2571	·2554	·2538	·2521	·2504	·2487	·2470	·2453	·2436	3	6	8	11	14
76	·2419	·2402	·2385	·2368	·2351	·2334	·2317	·2300	·2284	·2267	3	6	8	11	14
77	·2250	·2233	·2215	·2198	·2181	·2164	·2147	·2130	·2113	·2096	3	6	9	11	14
78	·2079	·2062	·2045	·2028	·2011	·1994	·1977	·1959	·1942	·1925	3	6	9	11	14
79	·1908	·1891	·1874	·1857	·1840	·1822	·1805	·1788	·1771	·1754	3	6	9	11	14
80	·1736	·1719	·1702	·1685	·1668	·1650	·1633	·1616	·1599	·1582	3	6	9	11	14
81	·1564	·1547	·1530	·1513	·1495	·1478	·1461	·1444	·1426	·1409	3	6	9	12	14
82	·1392	·1374	·1357	·1340	·1323	·1305	·1288	·1271	·1253	·1236	3	6	9	12	14
83	·1219	·1201	·1184	·1167	·1149	·1132	·1115	·1097	·1080	·1063	3	6	9	12	14
84	·1045	·1028	·1011	·993	·976	·958	·941	·924	·906	·889	3	6	9	12	14
85	·0872	·0854	·0837	·0819	·0802	·0785	·0767	·0750	·0732	·0715	3	6	9	12	14
86	·0698	·0680	·0663	·0645	·0628	·0610	·0593	·0576	·0558	·0541	3	6	9	12	15
87	·0523	·0506	·0488	·0471	·0454	·0436	·0419	·0401	·0384	·0366	3	6	9	12	15
88	·0349	·0332	·0314	·0297	·0279	·0262	·0244	·0227	·0209	·0192	3	6	9	12	15
89	·0175	·0157	·0140	·0122	·0105	·0087	·0070	·0052	·0035	·0017	3	6	9	12	15
	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	Subtract Differences.				
											1'	2'	3'	4'	5'

NATURAL TANGENTS

	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'
0°	'0000	'0017	'0035	'0052	'0070	'0087	'0105	'0122	'0140	'0157	3	6	9	12	15
1	'0175	0192	0209	0227	0244	0262	0279	0297	0314	0332	3	6	9	12	15
2	'0349	0367	0384	0402	0419	0437	0454	0472	0489	0507	3	6	9	12	15
3	'0524	0542	0559	0577	0594	0612	0629	0647	0664	0682	3	6	9	12	15
4	'0699	0717	0734	0752	0769	0787	0805	0822	0840	0857	3	6	9	12	15
5	'0875	0892	0910	0928	0945	0963	0981	0998	1016	1033	3	6	9	12	15
6	'1051	1069	1086	1104	1122	1139	1157	1175	1192	1210	3	6	9	12	15
7	'1228	1246	1263	1281	1299	1317	1334	1352	1370	1388	3	6	9	12	15
8	'1405	1423	1441	1459	1477	1495	1512	1530	1548	1566	3	6	9	12	15
9	'1584	1602	1620	1638	1655	1673	1691	1709	1727	1745	3	6	9	12	15
10	'1763	1781	1799	1817	1835	1853	1871	1890	1908	1926	3	6	9	12	15
11	'1944	1962	1980	1998	2016	2035	2053	2071	2089	2107	3	6	9	12	15
12	'2126	2144	2162	2180	2199	2217	2235	2254	2272	2290	3	6	9	12	15
13	'2309	2327	2345	2364	2382	2401	2419	2438	2456	2475	3	6	9	12	15
14	'2493	2512	2530	2549	2568	2586	2605	2623	2642	2661	3	6	9	12	10
15	'2679	2698	2717	2736	2754	2773	2792	2811	2830	2849	3	6	9	13	16
16	'2867	2886	2905	2924	2943	2962	2981	3000	3019	3038	3	6	9	13	16
17	'3057	3076	3096	3115	3134	3153	3172	3191	3211	3230	3	6	10	13	16
18	'3249	3269	3288	3307	3327	3346	3365	3385	3404	3424	3	6	10	13	16
19	'3443	3463	3482	3502	3522	3541	3561	3581	3600	3620	3	7	10	13	16
20	'3640	3659	3679	3699	3719	3739	3759	3779	3799	3819	3	7	10	13	17
21	'3839	3859	3879	3899	3919	3939	3959	3979	4000	4020	3	7	10	13	17
22	'4040	4061	4081	4101	4122	4142	4163	4183	4204	4224	3	7	10	14	17
23	'4245	4265	4286	4307	4327	4348	4369	4390	4411	4431	3	7	10	14	17
24	'4452	4473	4494	4515	4536	4557	4578	4599	4621	4642	4	7	11	14	18
25	'4663	4684	4706	4727	4748	4770	4791	4813	4834	4856	4	7	11	14	18
26	'4877	4899	4921	4942	4964	4986	5008	5029	5051	5073	4	7	11	15	18
27	'5095	5117	5139	5161	5184	5206	5228	5250	5272	5295	4	7	11	15	18
28	'5317	5340	5362	5384	5407	5430	5452	5475	5498	5520	4	8	11	15	19
29	'5543	5566	5589	5612	5635	5658	5681	5704	5727	5750	4	8	12	15	19
30	'5774	5797	5820	5844	5867	5890	5914	5938	5961	5985	4	8	12	16	20
31	'6009	6032	6056	6080	6104	6128	6152	6176	6200	6224	4	8	12	16	20
32	'6249	6273	6297	6322	6346	6371	6395	6420	6445	6469	4	8	12	16	20
33	'6494	6519	6544	6569	6594	6619	6644	6669	6694	6720	4	8	13	17	21
34	'6745	6771	6796	6822	6847	6873	6899	6924	6950	6976	4	9	13	17	21
35	'7002	7028	7054	7080	7107	7133	7159	7186	7212	7239	4	9	13	18	22
36	'7265	7292	7319	7346	7373	7400	7427	7454	7481	7508	5	9	14	18	23
37	'7536	7563	7590	7618	7646	7673	7701	7729	7757	7785	5	9	14	18	23
38	'7813	7841	7869	7898	7926	7954	7983	8012	8040	8069	5	9	14	19	24
39	'8098	8127	8156	8185	8214	8243	8273	8302	8332	8361	5	10	15	20	24
40	'8391	8421	8451	8481	8511	8541	8571	8601	8632	8662	5	10	15	20	25
41	'8693	8724	8754	8785	8816	8847	8878	8910	8941	8972	5	10	16	21	26
42	'9004	9036	9067	9099	9131	9163	9195	9228	9260	9293	5	11	16	21	27
43	'9325	9358	9391	9424	9457	9490	9523	9556	9590	9623	6	11	17	22	28
44	'9657	9691	9725	9759	9793	9827	9861	9896	9930	9965	6	11	17	23	29
	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'

	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1' 2'	3'	4'	5'
45°	1'0000	'0035	'0070	'0105	'0141	'0176	'0212	'0247	'0283	'0319	6 12	18	24	30
46	1'0355	0392	0428	0464	0501	0538	0575	0612	0649	0686	6 12	18	25	31
47	1'0724	0761	0799	0837	0875	0913	0951	0990	1028	1067	6 13	19	25	32
48	1'1106	1145	1184	1224	1263	1303	1343	1383	1423	1463	7 13	20	27	33
49	1'1504	1544	1585	1626	1667	1708	1750	1792	1833	1875	7 14	21	28	34
50	1'1918	1960	2002	2045	2088	2131	2174	2218	2261	2305	7 14	22	29	36
51	1'2349	2393	2437	2482	2527	2572	2617	2662	2708	2753	8 15	23	30	38
52	1'2799	2846	2892	2938	2985	3032	3079	3127	3175	3222	8 16	24	31	39
53	1'3270	3319	3367	3416	3465	3514	3564	3613	3663	3713	8 16	25	33	41
54	1'3764	3814	3865	3916	3968	4019	4071	4124	4176	4229	9 17	26	34	43
55	1'4281	4335	4388	4442	4496	4550	4605	4659	4715	4770	9 18	27	36	45
56	1'4826	4882	4938	4994	5051	5108	5166	5224	5282	5340	10 19	29	38	48
57	1'5399	5458	5517	5577	5637	5697	5757	5818	5880	5941	10 20	30	40	50
58	1'6003	6066	6128	6191	6255	6319	6383	6447	6512	6577	11 21	32	43	53
59	1'6643	6709	6775	6842	6909	6977	7045	7113	7182	7251	11 23	34	45	57
60	1'7321	7391	7461	7532	7603	7675	7747	7820	7893	7966	12 24	36	48	60
61	1'8040	8115	8190	8265	8341	8418	7495	8572	8650	8728	13 26	38	51	64
62	1'8807	8887	8967	9047	9128	9210	9292	9375	9458	9542	14 27	41	55	68
63	1'9626	9711	9797	9883	9970	2'0057	2'0145	2'0233	2'0323	2'0413	15 29	44	58	73
64	2'0503	0594	0686	0778	0872	0965	1060	1155	1251	1348	16 31	47	63	79
65	2'1445	1543	1642	1742	1842	1943	2045	2148	2251	2355	17 34	51	68	85
66	2'2460	2566	2673	2781	2889	2998	3109	3220	3332	3445	18 37	55	73	92
67	2'3559	3673	3789	3906	4023	4142	4262	4383	4504	4627	20 40	60	79	99
68	2'4751	4876	5002	5129	5257	5386	5517	5649	5782	5916	22 43	65	87	108
69	2'6051	6187	6325	6464	6605	6746	6889	7034	7179	7326	24 48	71	95	119
70	2'7475	7625	7776	7929	8083	8239	8397	8556	8716	8878	26 52	78	105	131
71	2'9042	9208	9375	9544	9714	9887	3'0061	3'0237	3'0415	3'0595	29 58	87	116	145
72	3'0777	0961	1146	1334	1524	1716	1910	2106	2305	2506	32 64	96	129	161
73	3'2709	2914	3122	3332	3544	3759	3977	4197	4420	4646	36 72	108	144	180
74	3'4874	5105	5339	5576	5816	6059	6305	6554	6806	7062	41 81	122	163	204
75	3'7321	7583	7848	8118	8391	8667	8947	9232	9520	9812	46 93	139	186	232
76	4'0108	0408	0713	1022	1335	1653	1976	2303	2635	2972	Mean differences no longer suffi- ciently accurate.			
77	4'3315	3662	4015	4374	4737	5107	5483	5864	6252	6646				
78	4'7046	7453	7867	8288	8716	9152	9594	5'0045	5'0504	5'0970				
79	5'1446	1929	2422	2924	3435	3955	4486	5026	5578	6140				
80	5'6713	7297	7894	8502	9124	9758	6'0405	6'1066	6'1742	6'2432				
81	6'3138	3859	4596	5350	6122	6912	7720	8548	9395	7'0264				
82	7'1154	2066	3002	3962	4947	5958	6996	8062	9158	8'0285				
83	8'1443	2636	3863	5126	6427	7769	9152	9'0579	9'2052	9'3572				
84	9'5114	9'677	9'845	10'02	10'20	10'39	10'58	10'78	10'99	11'20				
85	11'43	11'66	11'91	12'16	12'43	12'71	13'00	13'30	13'62	13'95				
86	14'30	14'67	15'06	15'46	15'89	16'35	16'83	17'34	17'89	18'46				
87	19'08	19'74	20'45	21'20	22'02	22'90	23'86	24'90	26'03	27'27				
88	28'64	30'14	31'82	33'69	35'80	38'19	40'92	44'07	47'74	52'08				
89	57'29	63'66	71'62	81'85	95'49	114'6	143'2	191'0	286'5	573'0				
	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'				

	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'
0°	0000	0017	0035	0052	0070	0087	0105	0122	0140	0157	3	6	9	12	15
1	0175	0192	0209	0227	0244	0262	0279	0297	0314	0332	3	6	9	12	15
2	0349	0367	0384	0401	0419	0436	0454	0471	0489	0506	3	6	9	12	15
3	0524	0541	0559	0576	0593	0611	0628	0646	0663	0681	3	6	9	12	15
4	0698	0716	0733	0750	0768	0785	0803	0820	0838	0855	3	6	9	12	15
5	0873	0890	0908	0925	0942	0960	0977	0995	1012	1030	3	6	9	12	15
6	1047	1065	1082	1100	1117	1134	1152	1169	1187	1204	3	6	9	12	15
7	1222	1239	1257	1274	1292	1309	1326	1344	1361	1379	3	6	9	12	15
8	1396	1414	1431	1449	1466	1484	1501	1518	1536	1553	3	6	9	12	15
9	1571	1588	1606	1623	1641	1658	1676	1693	1710	1728	3	6	9	12	15
10	1745	1763	1780	1798	1815	1833	1850	1868	1885	1902	3	6	9	12	15
11	1920	1937	1955	1972	1990	2007	2025	2042	2059	2077	3	6	9	12	15
12	2094	2112	2129	2147	2164	2182	2199	2217	2234	2251	3	6	9	12	15
13	2269	2286	2304	2321	2339	2356	2374	2391	2409	2426	3	6	9	12	15
14	2443	2461	2478	2496	2513	2531	2548	2566	2583	2601	3	6	9	12	15
15	2618	2635	2653	2670	2688	2705	2723	2740	2758	2775	3	6	9	12	15
16	2793	2810	2827	2845	2862	2880	2897	2915	2932	2950	3	6	9	12	15
17	2967	2985	3002	3019	3037	3054	3072	3089	3107	3124	3	6	9	12	15
18	3142	3159	3176	3194	3211	3229	3246	3264	3281	3299	3	6	9	12	15
19	3316	3334	3351	3368	3386	3403	3421	3438	3456	3473	3	6	9	12	15
20	3491	3508	3526	3543	3560	3578	3595	3613	3630	3648	3	6	9	12	15
21	3665	3683	3700	3718	3735	3752	3770	3787	3805	3822	3	6	9	12	15
22	3840	3857	3875	3892	3910	3927	3944	3962	3979	3997	3	6	9	12	15
23	4014	4032	4049	4067	4084	4102	4119	4136	4154	4171	3	6	9	12	15
24	4189	4206	4224	4241	4259	4276	4294	4311	4328	4346	3	6	9	12	15
25	4363	4381	4398	4416	4433	4451	4468	4485	4503	4520	3	6	9	12	15
26	4538	4555	4573	4590	4608	4625	4643	4660	4677	4695	3	6	9	12	15
27	4712	4730	4747	4765	4782	4800	4817	4835	4852	4869	3	6	9	12	15
28	4887	4904	4922	4939	4957	4974	4992	5009	5027	5044	3	6	9	12	15
29	5061	5079	5096	5114	5131	5149	5166	5184	5201	5219	3	6	9	12	15
30	5236	5253	5271	5288	5306	5323	5341	5358	5376	5393	3	6	9	12	15
31	5411	5428	5445	5463	5480	5498	5515	5533	5550	5568	3	6	9	12	15
32	5585	5603	5620	5637	5655	5672	5690	5707	5725	5742	3	6	9	12	15
33	5760	5777	5794	5812	5829	5847	5864	5882	5899	5917	3	6	9	12	15
34	5934	5952	5969	5986	6004	6021	6039	6056	6074	6091	3	6	9	12	15
35	6109	6126	6144	6161	6178	6196	6213	6231	6248	6266	3	6	9	12	15
36	6283	6301	6318	6336	6353	6370	6388	6405	6423	6440	3	6	9	12	15
37	6458	6475	6493	6510	6528	6545	6562	6580	6597	6615	3	6	9	12	15
38	6632	6650	6667	6685	6702	6720	6737	6754	6772	6789	3	6	9	12	15
39	6807	6824	6842	6859	6877	6894	6912	6929	6946	6964	3	6	9	12	15
40	6981	6999	7016	7034	7051	7069	7086	7103	7121	7138	3	6	9	12	15
41	7156	7173	7191	7208	7226	7243	7261	7278	7295	7313	3	6	9	12	15
42	7330	7348	7365	7383	7400	7418	7435	7453	7470	7487	3	6	9	12	15
43	7505	7522	7540	7557	7575	7592	7610	7627	7645	7662	3	6	9	12	15
44	7679	7697	7714	7732	7749	7767	7784	7802	7819	7837	3	6	9	12	15
	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'

	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'
45°	7854	7871	7889	7906	7924	7941	7959	7976	7994	8011	3	6	9	12	15
46	8029	8046	8063	8081	8098	8116	8133	8151	8168	8186	3	6	9	12	15
47	8203	8221	8238	8255	8273	8290	8308	8325	8343	8360	3	6	9	12	15
48	8378	8395	8412	8430	8447	8465	8482	8500	8517	8535	3	6	9	12	15
49	8552	8570	8587	8604	8622	8639	8657	8674	8692	8709	3	6	9	12	15
50	8727	8744	8762	8779	8796	8814	8831	8849	8866	8884	3	6	9	12	15
51	8901	8919	8936	8954	8971	8988	9006	9023	9041	9058	3	6	9	12	15
52	9076	9093	9111	9128	9146	9163	9180	9198	9215	9233	3	6	9	12	15
53	9250	9268	9285	9303	9320	9338	9355	9372	9390	9407	3	6	9	12	15
54	9425	9442	9460	9477	9495	9512	9529	9547	9564	9582	3	6	9	12	15
55	9599	9617	9634	9652	9669	9687	9704	9721	9739	9756	3	6	9	12	15
56	9774	9791	9809	9826	9844	9861	9879	9896	9913	9931	3	6	9	12	15
57	9948	9966	9983	10001	10018	10036	10053	10071	10088	10105	3	6	9	12	15
58	10123	0140	0158	0175	0193	0210	0228	0245	0263	0280	3	6	9	12	15
59	10297	0315	0332	0350	0367	0385	0402	0420	0437	0455	3	6	9	12	15
60	10472	0489	0507	0524	0542	0559	0577	0594	0612	0629	3	6	9	12	15
61	10647	0664	0681	0699	0716	0734	0751	0769	0786	0804	3	6	9	12	15
62	10821	0838	0856	0873	0891	0908	0926	0943	0961	0978	3	6	9	12	15
63	10996	1013	1030	1048	1065	1083	1100	1118	1135	1153	3	6	9	12	15
64	11170	1188	1205	1222	1240	1257	1275	1292	1310	1327	3	6	9	12	15
65	11345	1362	1380	1397	1414	1432	1449	1467	1484	1502	3	6	9	12	15
66	11519	1537	1554	1572	1589	1606	1624	1641	1659	1676	3	6	9	12	15
67	11694	1711	1729	1746	1764	1781	1798	1816	1833	1851	3	6	9	12	15
68	11868	1886	1903	1921	1938	1956	1973	1990	2008	2025	3	6	9	12	15
69	12043	2060	2078	2095	2113	2130	2147	2165	2182	2200	3	6	9	12	15
70	12217	2235	2252	2270	2287	2305	2322	2339	2357	2374	3	6	9	12	15
71	12392	2409	2427	2444	2462	2479	2497	2514	2531	2549	3	6	9	12	15
72	12566	2584	2601	2619	2636	2654	2671	2689	2706	2723	3	6	9	12	15
73	12741	2758	2776	2793	2811	2828	2846	2863	2881	2898	3	6	9	12	15
74	12915	2933	2950	2968	2985	3003	3020	3038	3055	3073	3	6	9	12	15
75	13090	3107	3125	3142	3160	3177	3195	3212	3230	3247	3	6	9	12	15
76	13265	3282	3299	3317	3334	3352	3369	3387	3404	3422	3	6	9	12	15
77	13439	3456	3474	3491	3509	3526	3544	3561	3579	3596	3	6	9	12	15
78	13614	3631	3648	3666	3683	3701	3718	3736	3753	3771	3	6	9	12	15
79	13788	3806	3823	3840	3858	3875	3893	3910	3928	3945	3	6	9	12	15
80	13963	3980	3998	4015	4032	4050	4067	4085	4102	4120	3	6	9	12	15
81	14137	4155	4172	4190	4207	4224	4242	4259	4277	4294	3	6	9	12	15
82	14312	4329	4347	4364	4382	4399	4416	4434	4451	4469	3	6	9	12	15
83	14486	4504	4521	4539	4556	4573	4591	4608	4626	4643	3	6	9	12	15
84	14661	4678	4696	4713	4731	4748	4765	4783	4800	4818	3	6	9	12	15
85	14835	4853	4870	4888	4905	4923	4940	4957	4975	4992	3	6	9	12	15
86	15010	5027	5045	5062	5080	5097	5115	5132	5149	5167	3	6	9	12	15
87	15184	5202	5219	5237	5254	5272	5289	5307	5324	5341	3	6	9	12	15
88	15359	5376	5394	5411	5429	5446	5464	5481	5499	5516	3	6	9	12	15
89	15533	5551	5568	5586	5603	5621	5638	5656	5673	5691	3	6	9	12	15
	0'	6'	12'	18'	24'	30'	36'	42'	48'	54'	1'	2'	3'	4'	5'

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